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(54) Title: COTTON FIBER TRANSCRIPTIONAL FACTORS

(57) Abstract

Novel DNA constructs are provided which may be used as molecular probes or inserted into a plant host to provide for modification of transcription of a DNA sequence of interest during various stages of cotton fiber development. The DNA constructs comprise a cotton fiber transcriptional initiation regulatory region associated with a gene which is expressed in cotton fiber. Also provided is novel cotton having a cotton fiber which has a natural color introduced by the expression in the cotton fiber cell, using such a construct, of pigment synthesis genes. Cotton fiber cells having color produced by genetic engineering and cotton cells comprising melanin and indigo pigments are included.

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## COTTON FIBER TRANSCRIPTIONAL FACTORS

CROSS-REFERENCE TO RELATED APPLICATIONS

5 This application is a continuation in part of United States application Serial No. 08/487,087 filed June 7, 1995, and a continuation in part of United States application Serial No. 08/480,178, filed June 7, 1995.

10

INTRODUCTIONTechnical Field

This invention relates to methods of using *in vitro* constructed DNA transcription or expression cassettes capable of directing fiber-tissue transcription of a DNA sequence of interest 15 in plants to produce fiber cells having an altered phenotype, and to methods of providing for or modifying various characteristics of cotton fiber. The invention is exemplified by methods of using cotton fiber promoters for altering the phenotype of cotton fiber, and cotton fibers produced by the method.

20

Background

In general, genetic engineering techniques have been directed to modifying the phenotype of individual prokaryotic and eukaryotic cells, especially in culture. Plant cells have proven 25 more intransigent than other eukaryotic cells, due not only to a lack of suitable vector systems but also as a result of the different goals involved. For many applications, it is desirable

to be able to control gene expression at a particular stage in the growth of a plant or in a particular plant part. For this purpose, regulatory sequences are required which afford the desired initiation of transcription in the appropriate cell types  
5 and/or at the appropriate time in the plant's development without having serious detrimental effects on plant development and productivity. It is therefore of interest to be able to isolate sequences which can be used to provide the desired regulation of transcription in a plant cell during the growing cycle of the host  
10 plant.

One aspect of this interest is the ability to change the phenotype of particular cell types, such as differentiated epidermal cells that originate in fiber tissue, i.e. cotton fiber cells, so as to provide for altered or improved aspects of the  
15 mature cell type. Cotton is a plant of great commercial significance. In addition to the use of cotton fiber in the production of textiles, other uses of cotton include food preparation with cotton seed oil and animal feed derived from cotton seed husks.

20 Despite the importance of cotton as a crop, the breeding and genetic engineering of cotton fiber phenotypes has taken place at a relatively slow rate because of the absence of reliable promoters for use in selectively effecting changes in the phenotype of the fiber. In order to effect the desired phenotypic  
25 changes, transcription initiation regions capable of initiating transcription in fiber cells during development are desired. Thus, an important goal of cotton bioengineering research is the

acquisition of a reliable promoter which would permit expression of a protein selectively in cotton fiber to affect such qualities as fiber strength, length, color and dyability.

5 Relevant Literature

Cotton fiber-specific promoters are discussed in PCT publications WO 94/12014 and WO 95/08914, and John and Crow, Proc. Natl. Acad. Sci. USA, 89:5769-5773, 1992. cDNA clones that are preferentially expressed in cotton fiber have been isolated. One 10 of the clones isolated corresponds to mRNA and protein that are highest during the late primary cell wall and early secondary cell wall synthesis stages. John and Crow, *supra*.

In animals, the *ras* superfamily is subdivided into the subfamilies *ras* which is involved in controlling cell growth and division, *rab/YPT* members which control secretory processes, and *rho* which is involved in control of cytoskeletal organization (Bourne et al., (1991) Nature 349: 117-127), and number of homologous genes have now been identified in plants (for a review, see Terryn et al., (1993) Plant Mol. Biol. 22: 143-152). None 20 have been found for the important *ras* subfamily, all but one of the genes identified have been members of the *rab/YPT1* subfamily, and there is only one recent report of the cloning of a *rho* gene in pea (Yang and Watson(1993) Proc. Natl. Acad. Sci. USA 90: 8732-8736).

25 Little work has been done to characterize the functions of these genes in plants, although one recent report has shown that a small G protein from *Arabidopsis* can functionally complement a

mutant form in yeast involved in vesicle trafficking, suggesting a similar function for the plant gene (Bednarek et al., (1994) Plant Physiol 104: 591-596).

In animals, two members of the rho subfamily, called Rac and 5 Rho, have been shown to be involved in the regulation of actin organization (for a review, see Downward, (1992) Nature 359: 273-274).

Rac1 has been shown to mediate growth factor-induced membrane ruffling by influencing microfilament alignment on the plasma 10 membrane (Ridley et al, (1992) Cell 70: 401-410), whereas RhoA regulates the formation of actin stress fibers associated with focal adhesions (Ridley and Hall, (1992) Cell 70: 389-399).

In yeast, the CDC42 gene codes for a rho-type protein which also regulates actin organization involved in the establishment of 15 cell polarity required for the localized deposition of chitin in the bud scar (Adams et al., (1990) J Cell Biol 111: 131-143.

Disruption of gene function, either by temperature shifts with a CDC42-temperature-sensitive mutant in yeast (Adams et al., 1990), or by micro-injection into fibroblasts of mutant Rac or Rho 20 proteins exhibiting a dominant negative phenotype (Ridley et al., 1992; Ridley and Hall, 1992), leads to disorganization of the actin network.

In plants, control of cytoskeletal organization is poorly understood in spite of its importance for the regulation of 25 patterns of cell division, expansion, and subsequent deposition of secondary cell wall polymers. The cotton fiber represents an excellent system for studying cytoskeletal organization. Cotton

fibers are single cells in which cell elongation and secondary wall deposition can be studied as distinct events. These fibers develop synchronously within the boll following anthesis, and each fiber cell elongates for about 3 weeks, depositing a thin primary wall (Meinert and Delmer, (1984) Plant Physiol. 59: 1088-1097; Basra and Malik, (1984) Int Rev of Cytol 89: 65-113). At the time of transition to secondary wall cellulose synthesis, the fiber cells undergo a synchronous shift in the pattern of cortical microtubule and cell wall microfibril alignments, events which may 10 be regulated upstream by the organization of actin (Seagull, (1990) Protoplasma 159: 44-59; and (1992) In: Proceedings of the Cotton Fiber Cellulose Conference, National Cotton Council of America, Memphis TN, pp 171-192.

Agrobacterium-mediated cotton transformation is described in 15 Umbeck, United States Patents Nos. 5,004,863 and 5,159,135 and cotton transformation by particle bombardment is reported in WO 92/15675, published September 17, 1992. Transformation of *Brassica* has been described by Radke et al. (Theor. Appl. Genet. (1988) 75:685-694; Plant Cell Reports (1992) 11:499-505.

20

#### SUMMARY OF THE INVENTION

Novel DNA constructs and methods for their use are described which are capable of directing transcription of a gene of interest in cotton fiber, particularly early in fiber development and 25 during secondary cell wall development. The novel constructs include a vector comprising a transcriptional and translational initiation region obtainable from a gene expressed in cotton fiber

and methods of using constructs including the vector for altering fiber phenotype. Both the endogenous 3' regions and 5' regions may be important in directing efficient transcription and translation.

5       Three promoters are provided from genes involved in the regulation of cotton fiber development. One, Rac13, is from a protein in cotton which codes for an animal Rac protein homolog. Rac13, shows highly-enhanced expression during fiber development. This pattern of expression correlates well with the timing of  
10      reorganization of the cytoskeleton, suggesting that the Rac13 cotton gene may, like its animal counterpart, be involved in the signal transduction pathway for cytoskeletal organization. Rac13 is a gene that is moderately expressed during fiber development turning on at 9 dpa and shutting down approximately 24 dpa. It is  
15      maximally expressed between 17-21 dpa developing fiber.

Another promoter from a cotton protein is designated 4-4. The 4-4 mRNA accumulates in fiber cells at day 17 post anthesis and continues towards fiber maturity, which occurs at 60 days or so post anthesis. Data demonstrates that the 4-4 promoter remains  
20      very active at day 35 post anthesis.

Also provided is a promoter from a lipid transfer protein (hereinafter sometimes referred to as "Ltp") which is preferentially expressed in cotton fiber.

The methods of the present invention include transfecting a  
25      host plant cell of interest with a transcription or expression cassette comprising a cotton fiber promoter and generating a plant which is grown to produce fiber having the desired phenotype.

Constructs and methods of the subject invention thus find use in modulation of endogenous fiber products, as well as production of exogenous products and in modifying the phenotype of fiber and fiber products. The constructs also find use as molecular probes.

- 5 In particular, constructs and methods for use in gene expression in cotton embryo tissues are considered herein. By these methods, novel cotton plants and cotton plant parts, such as modified cotton fibers, may be obtained.

Also provided are constructs and methods of use relating to  
10 modification of color phenotype in cotton fiber. Such constructs contain sequences for expression of genes involved in the production of colored compounds, such as anthocyanins, melanin or indigo, and also may contain sequences which provide for targeting of the gene products to particular locations in the plant cell,  
15 such as plastid organelles, or vacuoles. Plastid targeting is of particular interest for expression of genes involved in aromatic amino acid biosynthesis pathways, while vacuolar targeting is of particular interest where the precursors required in synthesis of the pigment are present in vacuoles.

20 Of particular interest are plants producing fibers which are color, that is, with pigment produced in the fiber by the plant during fiber development, as opposed to fibers which are harvested and dyed or otherwise pigmented by separate processing. Fibers from a plant producing such colored fiber may be used to produce  
25 colored yarns and/or fabric which have not been subjected to any dyeing process. While naturally colored cotton has been available from various domesticated and wild type cotton varieties, th

instant application provides cotton fiber has a color produced by the expression of a genetically engineered protein.

Thus, the application provides constructs and methods of use relating to modification of color phenotype in cotton fiber. Such constructs contain sequences for expression of genes involved in the production of colored compounds, such as melanin or indigo, and also contain sequences which provide for targeting of the gene products to particular locations in the plant cell, such as plastid organelles, or vacuoles. Plastid targeting is of particular interest for expression of genes involved in the aromatic amino acid biosynthesis pathways, while vacuolar targeting is of particular interest where the precursors required in synthesis of the pigment are present in vacuoles.

15

#### DESCRIPTION OF THE DRAWINGS

Figure 1 shows the DNA sequence encoding the structural protein from cDNA 4-4.

Figure 2 shows the sequence to the promoter construct pCGN5606 made using genomic DNA from 4-4-6 genomic clone.

20 Figure 3 shows the sequence to the 4-4 promoter construct pCGN5610.

Figure 4 shows the cDNA sequence encoding the Rac13 gene expressed in cotton fiber.

25 Figure 5 shows the sequence the promoter region from the rac13 gene.

Figure 6 shows a restriction map for pCGN4735.

Figure 7 shows the sequence of the Ltp promoter region from a cotton fiber specific lipid transfer protein gene.

Figure 8 shows the arrangement of a binary vectors pCGN5148 and pCGN5616 for plant transformation to express genes for melanin synthesis and indigo synthesis, respectively.

Figure 9 provides the results of color measurements taken from fibers of the control Coker 130 cotton used in transformation using color constructs.

Figure 10 shows the results of measurements made of color of plants transformed by the pCGN5148 construct to express genes for melanin synthesis.

Figure 11 shows the results of measurements taken of the color of plants transformed by the pCGN5149 construct to express genes for melanin synthesis.

Figure 12 shows the results of measurements made of color of plants transformed to express genes for indigo synthesis, using construct pCGN5616.

Figure 13 shows control measurements made of naturally colored cotton plants which are produced by non-transgenic colored cotton plants.

#### DETAILED DESCRIPTION OF THE INVENTION

In accordance with the subject invention, novel constructs and methods are described, which may be used provide for transcription of a nucleotide sequence of interest in cells of a plant host, preferentially in cotton fiber cells to produce cotton fiber having an altered color phenotype.

Cotton fiber is a differentiated single epidermal cell of the outer integument of the ovule. It has four distinct growth phases; initiation, elongation (primary cell wall synthesis), secondary cell wall synthesis, and maturation. Initiation of 5 fiber development appears to be triggered by hormones. The primary cell wall is laid down during the elongation phase, lasting up to 25 days postanthesis (DPA). Synthesis of the secondary wall commences prior to the cessation of the elongation phase and continues to approximately 40 DPA, forming a wall of 10 almost pure cellulose.

The constructs for use in such cells may include several forms, depending upon the intended use of the construct. Thus, the constructs include vectors, transcriptional cassettes, expression cassettes and plasmids. The transcriptional and 15 translational initiation region (also sometimes referred to as a "promoter,"), preferably comprises a transcriptional initiation regulatory region and a translational initiation regulatory region of untranslated 5' sequences, "ribosome binding sites," responsible for binding mRNA to ribosomes and translational 20 initiation. It is preferred that all of the transcriptional and translational functional elements of the initiation control region are derived from or obtainable from the same gene. In some embodiments, the promoter will be modified by the addition of 25 sequences, such as enhancers, or deletions of nonessential and/or undesired sequences. By "obtainable" is intended a promoter having a DNA sequence sufficiently similar to that of a native promoter to provide for the desired specificity of transcription.

of a DNA sequence of interest. It includes natural and synthetic sequences as well as sequences which may be a combination of synthetic and natural sequences.

Cotton fiber transcriptional initiation regions chosen for

- 5 cotton fiber modification may include the 4-4, rac13 and Ltp  
cotton fiber promoter regions provided herein.

A transcriptional cassette for transcription of a nucleotide sequence of interest in cotton fiber will include in the direction of transcription, the cotton fiber transcriptional initiation 10 region, a DNA sequence of interest, and a transcriptional termination region functional in the plant cell. When the cassette provides for the transcription and translation of a DNA sequence of interest it is considered an expression cassette. One or more introns may be also be present.

- 15 Other sequences may also be present, including those encoding transit peptides and secretory leader sequences as desired.

Fiber-tissue transcription initiation regions of this invention are, preferably, not readily detectable in other plant tissues. Transcription initiation regions capable of initiating 20 transcription in other plant tissues and/or at other stages of fiber development, in addition to the foregoing, are acceptable insofar as such regions provide a significant expression level in cotton fiber at the defined periods of interest and do not negatively interfere with the plant as a whole, and, in 25 particular, do not interfere with the development of fiber and/or fiber-related parts.

Downstream from, and under the regulatory control of, the cotton fiber transcriptional/translational initiation control region is a nucleotide sequence of interest which provides for modification of the phenotype of fiber. The nucleotide sequence 5 may be any open reading frame encoding a polypeptide of interest, for example, an enzyme, or a sequence complementary to a genomic sequence, where the genomic sequence may be an open reading frame, an intron, a noncoding leader sequence, or any other sequence where the complementary sequence inhibits transcription, messenger 10 RNA processing, for example, splicing, or translation. The nucleotide sequences of this invention may be synthetic, naturally derived, or combinations thereof. Depending upon the nature of the DNA sequence of interest, it may be desirable to synthesize the sequence with plant preferred codons. The plant preferred 15 codons may be determined from the codons of highest frequency in the proteins expressed in the largest amount in the particular plant species of interest. Phenotypic modification can be achieved by modulating production either of an endogenous transcription or translation product, for example as to the 20 amount, relative distribution, or the like, or an exogenous transcription or translation product, for example to provide for a novel function or products in a transgenic host cell or tissue. Of particular interest are DNA sequences encoding expression 25 products associated with the development of plant fiber, including genes involved in metabolism of cytokinins, auxins, ethylene, abscisic acid, and the like. Methods and compositions for modulating cytokinin expression are described in United States

Patent No. 5,177,307, which disclosure is hereby incorporated by reference. Alternatively, various genes, from sources including other eukaryotic or prokaryotic cells, including bacteria, such as those from *Agrobacterium tumefaciens* T-DNA auxin and cytokinin biosynthetic gene products, for example, and mammals, for example interferons, may be used.

Other phenotypic modifications include modification of the color of cotton fibers. Of interest are genes involved in production of melanin and genes involved in the production of indigo. Melanins are dark brown pigments found in animals, plants and microorganisms, any of which may serve as a source for sequences for insertion into the constructs of the present invention. Specific examples include the tyrosinase gene which can be cloned from *Streptomyces antibioticus*. The ORF438 encoded protein in *S. antibioticus* also is necessary for melanin production, and may provide a copper donor function. In addition, a tyrosinase gene can be isolated from any organism which makes melanin. The gene can be isolated from human hair, melanocytes or melanomas, cuttle fish and red roosters, among others. See, for example, EP Application No. 89118346.9 which discloses a process for producing melanins, their precursors and derivatives in microorganisms. Also, See, Bernan et al. Gene (1985) 37:101-110; and della-Cioppa et al. Bio/Technology (1990) 8:634-638.

Indigo may be obtained by use of genes encoding a monooxygenase such as xylene oxygenase which oxidizes toluene and xylene to (methyl) benzyl alcohol and also transforms indole to indigo. Cloning of the xylene oxygenase gene and the nucleotide

and amino acid sequences are described in unexamined Japan se Patent Application Kokai:2-119777, published May 7, 1990. A dioxygenase such as naphthalene dioxygenase which also converts indole to indigo finds use; the naphthalene dioxygenase gene nahA 5 is described in Science (1983) 222: 167. For cloning, nucleotide sequence in characterization of genes encoding naphthalene dioxygenase of *Pseudomonas putida*. See, Kurkela et al. Gene (1988) 73:355-362. A tryptophanase gene sequence can be used in conjunction with an oxygenase to increase the amount of indole 10 available for conversion to indigo. Sources of tryptophanase gene sequences include *E. coli* (see, for example, Deeley et al. (1982) *J. Bacteriol.* 151 :942-951).

Plastid targeting sequences (transit peptides) are available from a number of plant nuclear-encoded plastid proteins, such as 15 the small subunit (SSU) of ribulose bisphosphate carboxylase, plant fatty acid biosynthesis related genes including acyl carrier protein (ACP), stearoyl-ACP desaturase,  $\beta$ -ketoacyl-ACP synthase and acyl-ACP thioesterase, or LHCPII genes. The encoding sequence for a transit peptide which provides for transport to plastids may 20 include all or a portion of the encoding sequence for a particular transit peptide, and may also contain portions of the mature protein encoding sequence associated with a particular transit peptide. There are numerous examples in the art of transit peptides which may be used to deliver a target protein into a 25 plastid organelle. The particular transit peptide encoding sequence used in the instant invention is not critical, as long as delivery to the plastid is obtained.

As an alternative to using transit peptides to target pigment synthesis proteins to plastid organelles, the desired constructs may be used to transform the plastid genome directly. In this instance, promoters capable of providing for transcription of genes in plant plastids are desired. Of particular interest is the use of a T7 promoter to provide for high levels of transcription. Since plastids do not contain an appropriate polymerase for transcription from the T7 promoter, T7 polymerase may be expressed from a nuclear construct and targeted to plastids using transit peptides as described above. (See McBride et al. (1994) *Proc. Nat. Acad. Sci.* 91:7301-7305; see also copending US patent application entitled "Controlled Expression of Transgenic Constructs in Plant Plastids", serial no. 08/472,719, filed June 6, 1995, and copending US patent application SN 08/167,638, filed December 14, 1993 and PCT/US94/14574 filed December 12, 1994.) Tissue specific or developmentally regulated promoters may be useful for expression of the T7 polymerase in order to limit expression to the appropriate tissue or stage of development.

Targeting of melanin synthesis genes to vacuoles is also of interest in plant tissues which accumulate the tyrosine substrate involved in melanin synthesis in vacuoles. The protein signal for targeting to vacuoles may be provided from a plant gene which is normally transported across the rough endoplasmic reticulum, such as the 32 amino acid N-terminal region of the metallocarboxypeptidase inhibitor gene from tomato (Martineau et al. (1991) *Mol. Gen. Genet.* 228 :281-286). In addition to the signal sequence, vacuolar targeting constructs also encode a

vacuolar localization signal (VLS) positioned at the carboxy terminus of the encoded protein. Appropriate signal sequences and VLS regions may be obtained from various other plant genes and may be similarly used in the constructs of this invention. Numerous 5 vacuolar targetting peptides are known to the art, as are reviewed in Chrispeels et al., Cell (1992) 68:613-616.

The Maize A1 gene which encodes a dihydroflavonol reductase, an enzyme of the anthocyanin pigmentation pathway is one such gene. In cells that express the A1 gene, dihydrokempferol is 10 converted to 2-8 alkylleucopelargonidin, which may be further metabolized to pelargonidin pigment by endogenous plant enzymes. Other anthocyanin or flavonoid type pigments may also be of interest for modification of cotton cell fibers, and have been suggested for use in plant flowers (for a review of plant flower 15 color, see van Tunen et al., Plant Biotechnology Series, Volume 2 (1990) Developmental Regulation of Plant Gene Expression, D. Grierson ed.). Anthocyanin is produced by a progression of steps from cellular phenylalanine pools. The R and C1 genes are maize regulatory proteins which are active by positively affecting 20 upstream steps in the anthocyanin biosynthesis from these pools. The R gene is described in Perot and Cone (1989) Nucl. Acids Res., 17:8003, and the C1 gene is described in Paz-Ares et al (1987) EMBO, 6:3553-3558. Lloyd et al. (1992) Science, 258:1773-1775 discussed both genes.

25 Although cotton fibers in commercially grown varieties are primarily white in color, other naturally occurring cotton varieties have brown or reddish-brown fibers. Additionally, a

cotton line containing green colored fibers has been identified. Cotton lines providing such fibers are available from various sources, including the BC variety cottons (BC Cotton Inc., Box 8656, Bakersfield, CA 93389) and Fox Fibre cottons (Natural 5 Cotton Colors, Inc., P.O. Box 791, Wasco, CA 93280).

The existence of such colored cotton lines suggests that the precursors required for the anthocyanin pigment pathways are present in cotton fibers cells, thus allowing further color phenotype modifications. Thus, the maize R and C1 genes could be 10 used in enhancing the levels of of anthocyanin produced in fiber cells. As the R and C1 proteins are proteins with a positive control at the regulatory level on anthocyanin pigment precursor biosynthesis, these proteins are expressed in the nucleus, and not targetted to plastids or vacuoles.

15 For some applications, it is of interest to modify other aspects of the fiber. For example, it is of interest to modify various aspects of cotton fibers, such as strength or texture of a fiber. Thus, the appropriate gene may be inserted in the constructs of the invention, including genes for PHB biosynthesis 20 (see, Peoples et al. *J. Biol. Chem.* (1989) 264: 15298-15303 and *Ibid.* 15293-15397; Saxena, *Plant Molecular Biology* (1990) 15:673-683, which discloses cloning and sequencing of the cellulose synthase catalytic subunit gene; and Bowen et al. *PNAS* (1992) 89:519-523 which discloses chitin synthase genes of *Saccharomyces* 25 *cerevisiae* and *Candida albicans*. Various constructs and methods are disclosed for the use of hormones to effect changes to fiber quality in copending US patent application entitled "Cotton

"Modification Using Ovary-Tissue Transcriptional factors", serial no. 08/397,652 filed February 2, 1995, the teachings of which are incorporated herein by reference.

Transcriptional cassettes may be used when the transcription 5 of an anti-sense sequence is desired. When the expression of a polypeptide is desired, expression cassettes providing for transcription and translation of the DNA sequence of interest will be used. Various changes are of interest; these changes may include modulation (increase or decrease) of formation of 10 particular saccharides, hormones, enzymes, or other biological parameters. These also include modifying the composition of the final fiber that is changing the ratio and/or amounts of water, solids, fiber or sugars. Other phenotypic properties of interest for modification include response to stress, organisms, 15 herbicides, brushing, growth regulators, and the like. These results can be achieved by providing for reduction of expression of one or more endogenous products, particularly an enzyme or cofactor, either by producing a transcription product which is complementary (anti-sense) to the transcription product of a 20 native gene, so as to inhibit the maturation and/or expression of the transcription product, or by providing for expression of a gene, either endogenous or exogenous, to be associated with the development of a plant fiber.

The termination region which is employed in the expression 25 cassette will be primarily one of convenience, since the termination regions appear to be relatively interchangeable. The termination region may be native with the transcriptional

initiation region, may be native with the DNA sequence of interest, may be derived from another source. The termination region may be naturally occurring, or wholly or partially synthetic. Convenient termination regions are available from the 5 Ti-plasmid of *A. tumefaciens*, such as the octopine synthase and nopaline synthase termination regions. In some embodiments, it may be desired to use the 3' termination region native to the cotton fiber transcription initiation region used in a particular construct.

10 As described herein, in some instances additional nucleotide sequences will be present in the constructs to provide for targeting of a particular gene product to specific cellular locations. For example, where coding sequences for synthesis of aromatic colored pigments are used in a construct, particularly 15 coding sequences for enzymes which have as their substrates aromatic compounds such tyrosine and indole, it is preferable to include sequences which provide for delivery of the enzyme into plastids, such as an SSU transit peptide sequence. Also, for synthesis of pigments derived from tyrosine, such as melanin, 20 targeting to the vacuole may provide for enhanced color modifications.

For melanin production, the tyrosinase and ORF438 genes from *Streptomyces antibioticus* (Berman et al. (1985) 37:101-110) are provided in cotton fiber cells for expression from a 4-4 and Rac13 25 promoter. In *Streptomyces*, the ORF438 and tyrosinase proteins are expressed from the same promoter region. For expression from constructs in a transgenic plant genome, the coding regions may be

provided under the regulatory control of separate promoter regions. The promoter regions may be the same or different for the two genes. Alternatively, coordinate expression of the two genes from a single plant promoter may be desired. Constructs for 5 expression of the tyrosinase and ORF438 gene products from 4-4 and rac promoter regions are described in detail in the following examples. Additional promoters may also be desired, for example plant viral promoters, such as CaMV 35S, can be used for constitutive expression of one of the desired gene products, with 10 the other gene product being expressed in cotton fiber tissues from the 4-4 and rac promoter.

Similarly, other constitutive promoters may also be useful in certain applications, for example the mas, Mac or DoubleMac, promoters described in United States Patent No. 5,106,739 and by 15 Comai et al., *Plant Mol. Biol.* (1990) 15:373-381). When plants comprising multiple gene constructs are desired, for example plants expressing the melanin genes, ORF438 and tyrosinase, the plants may be obtained by co-transformation with both constructs, or by transformation with individual constructs followed by plant 20 breeding methods to obtain plants expressing both of the desired genes.

A variety of techniques are available and known to those skilled in the art for introduction of constructs into a plant cell host. These techniques include transfection with DNA 25 employing *A. tumefaciens* or *A. rhizogenes* as the transfecting agent, protoplast fusion, injection, electroporation, particle acceleration, etc. For transformation with *Agrobacterium*,

plasmids can be prepared in *E. coli* which contain DNA homologous with the Ti-plasmid, particularly T-DNA. The plasmid may or may not be capable of replication in *Agrobacterium*, that is, it may or may not have a broad spectrum prokaryotic replication system such as does, for example, pRK290, depending in part upon whether the transcription cassette is to be integrated into the Ti-plasmid or to be retained on an independent plasmid. The *Agrobacterium* host will contain a plasmid having the *vir* genes necessary for transfer of the T-DNA to the plant cell and may or may not have the complete T-DNA. At least the right border and frequently both the right and left borders of the T-DNA of the Ti- or Ri-plasmids will be joined as flanking regions to the transcription construct. The use of T-DNA for transformation of plant cells has received extensive study and is amply described in EPA Serial No. 120,516, Hoekema, In: The Binary Plant Vector System Offset-drukkerij Kanters B.V., Albllasserdam, 1985, Chapter V, Knauf, et al., Genetic Analysis of Host Range Expression by *Agrobacterium*, In: Molecular Genetics of the Bacteria-Plant Interaction, Puhler, A. ed., Springer-Verlag, NY, 1983, p. 245, and An, et al., EMBO J. (1985) 4:277-284.

For infection, particle acceleration and electroporation, a disarmed Ti-plasmid lacking particularly the tumor genes found in the T-DNA region) may be introduced into the plant cell. By means of a helper plasmid, the construct may be transferred to the *A. tumefaciens* and the resulting transfected organism used for transfecting a plant cell; explants may be cultivated with transformed *A. tumefaciens* or *A. rhizogenes* to allow for transfer

of the transcription cassette to the plant cells. Alternatively, to enhance integration into the plant genome, terminal repeats of transposons may be used as borders in conjunction with a transposase. In this situation, expression of the transposase 5 should be inducible, so that once the transcription construct is integrated into the genome, it should be relatively stably integrated. Transgenic plant cells are then placed in an appropriate selective medium for selection of transgenic cells which are then grown to callus, shoots grown and plantlets 10 generated from the shoot by growing in rooting medium.

To confirm the presence of the transgenes in transgenic cells and plants, a Southern blot analysis can be performed using methods known to those skilled in the art. Expression products of the transgenes can be detected in any of a variety of ways, 15 depending upon the nature of the product, and include immune assay, enzyme assay or visual inspection, for example to detect pigment formation in the appropriate plant part or cells. Once transgenic plants have been obtained, they may be grown to produce fiber having the desired phenotype. The fibers may be harvested, 20 and/or the seed collected. The seed may serve as a source for growing additional plants having the desired characteristics. The terms transgenic plants and transgenic cells include plants and cells derived from either transgenic plants or transgenic cells.

The various sequences provided herein may be used as 25 molecular probes for the isolation of other sequences which may be useful in the present invention, for example, to obtain related transcriptional initiation regions from the same or different

plant sources. Related transcriptional initiation regions obtainable from the sequences provided in this invention will show at least about 60% homology, and more preferred regions will demonstrate an even greater percentage of homology with the 5 probes. Of particular importance is the ability to obtain related transcription initiation control regions having the timing and tissue parameters described herein. For example, using the probe 4-4 and rac, at least 7 additional clones, have been identified, but not further characterized. Thus, by employing the techniques 10 described in this application, and other techniques known in the art (such as Maniatis, et al., *Molecular Cloning, - A Laboratory Manual* (Cold Spring Harbor, New York) 1982), other transcription initiation regions capable of directing cotton fiber transcription as described in this invention may be determined. The constructs 15 can also be used in conjunction with plant regeneration systems to obtain plant cells and plants; thus, the constructs may be used to modify the phenotype of fiber cells, to provide cotton fibers which are colored as the result of genetic engineering to heretofor unavailable hues and/or intensities.

20 Various varieties and lines of cotton may find use in the described methods. Cultivated cotton species include *Gossypium hirsutum* and *G. babadense* (extra-long staple, or Pima cotton), which evolved in the New World, and the Old World crops *G. herbaceum* and *G. arboreum*.

25 Color phenotypes can be assessed by the use of a colorimeter, an instrument which is already used to provide objective measurements of the color of cotton samples. A colorimeter uses a

combination of light sources and filters to make various estimates of a samples colors, sometimes referred to as tristimulus values.

In the past such estimtes have been used to calculate a value (Hunter's + b, described below) indicating the degree of 5 yellowness of a cotton sample. The yellowness and reflectance (from Rd, the degree of lightness or darkness of the samples) has been used to provide cotton color measurements for grading. Tests are typically conducted by exposing the face of a sample to a controlled light source. A typical color chart showing how the 10 official grade standards relate to Rd and+ b measurements is shown in Cotton, RJ Kohel and CF Lewis, Editors #24 in AGRONOMY Series-American Soc. Agromony (see Fig. 12-6).

Various colorimeter methods can be so used to quantify color and express it numerically. The Munsell method, devised by the 15 American artist A.. Munsell, uses a classification system of paper color chips assorted according to their hue (Munsell Hue), lightness (Munsell Value), and saturation (Munsell Chroma) for visual comparison with a specimen color.

Other methods for expressing color numerically have been 20 developed by an international organization concerned with light and color, the Commission Internationale de l'Eclairage (CIE), having a Central Bureau located at Kegelgasse 27, A-1030 Vienna, AUSTRIA. The two most widely known of these methods are the Yxy color space, devised in 1931 based on the tristimulus value XYZ, 25 as defined by CIE, and the L\*a\*b\* color space, devised in 1976 to provide more uniform color differences in relation to visual differences. Color spaces\* such as these are now used throughout

the world for color communication. The Hunter Lab color space was developed in 1948 by R.S. Hunter as a uniform color space which could be read directly from a photoelectric colorimeter (tristimulus method).

5       The L\*C\*h color space uses the same diagram as the L\*a\*b\* color space, but uses cylindrical coordinates instead of rectangular coordinates. In this color space, L\* indicates lightness and is the same as the L\* of the L\*a\*b\* color space, C\* is chroma, and h is the hue angle. The value of chroma C is 0 at  
10      the center and increases according to the distance from the center. Hue angle is defined as starting at the +a axis of the L\*a\*b\* space, and is expressed in degrees in a counterclockwise rotation. Thus, relative to the L\*a\*b\* space, 0° and 360° would be at the +a\* line, 90° would be +b\*, 180° would be -a\* and 270°  
15      would be -b\*.

All of the above methods can be used to obtain precise measurements of a cotton fiber color phenotype.

#### EXPERIMENTAL

20       The following examples are offered by way of illustration and not by limitation.

##### Example 1

###### cDNA libraries

###### Tissue preparation for cDNA synthesis

25       Leaf and root tissue were isolated from 8 inch tall greenhouse grown seedlings and immediately frozen in liquid nitrogen. Flowers were collected at the rapidly expanding 3 day

- preatthesis stage and also frozen. Seed was collected from 21 day postanthesis locules which had been removed from the boll and frozen entire in liquid nitrogen. Once frozen, the fiber was removed from the seed and the denuded seed used for RNA isolation.
- 5 All fibers were removed from the seed under liquid nitrogen and the fiber was ground to a powder prior to RNA isolation. Fibers were from bolls which had been tagged at anthesis.

#### DNA and RNA Manipulations

- 10 The lambda ZapII™ cDNA library system of Stratagene was used for screening, and was prepared from cDNA derived from poly-A<sup>+</sup> mRNA isolated from fibers of *Gossypium hirsutum* cultivar Acala SJ-2. The fibers were isolated from bolls harvested at approximately 21 dpa using field-grown plants in Israel.
- 15 Total RNA was isolated from 21 dpa seeds (*G. hirsutum* cv Coker 130 from which the fiber had been removed) using the method of Hughes and Galau ((1988) Plant Mol Biol Reporter, 6:253-257.) All other RNAs were prepared according to Hall et al. ((1978), Proc Natl Acad Sci USA 75: 3196-3200), with the following
- 20 modifications. After the second 2M LiCl wash, the pellet was dissolved in 1/10 original volume of 10 mM Tris pH7.5 and brought to 35mM potassium acetate pH6.5 and 1/2 volume EtOH was added slowly. The mixture was placed on ice for 15 minutes and then centrifuged at 20,000 x g for 15 minutes at 4<sup>0</sup>C. The potassium
- 25 acetate concentration was brought to 0.2M, 2 1/2 volumes EtOH added and the RNA placed at -20<sup>0</sup>C for several hours. The precipitate was centrifuged at 12,000 x g for 30 minutes at 4<sup>0</sup>C

and the pellet was resuspended in diethylpyrocarbonate-treated water. Poly-A<sup>+</sup> RNA was prepared from total mRNA utilizing an oligo(dT)-cellulose kit (Becton Dickenson) and following the manufacturer's protocol.

5       Cotton genomic DNA was prepared as follows. Four grams of young cotton leaf tissue (cv Coker 130) was ground to a powder in N<sub>2</sub> and placed in an Oak Ridge tube with 0.4g polyvinylpyrrolidone and 20mls extraction buffer (200mM Ches/NaOH pH9.1, 200mM NaCl, 100mMEDTA/NaOH pH9.0, 2% SDS, 0.5% Na deoxycholate, 2% Nonidet NP-10, 20mM B-mercaptoethanol) was added to sample, gently mixed and incubated at 65<sup>0</sup>C in a shaking water bath for 10 minutes. 7.0 mls of 5M potassium acetate pH6.5 was added and carefully mixed. Incubation was carried out on ice for 30 minutes with gentle mixing every 5 minutes. The sample was centrifuged for 20 minutes 15 at 21,000 x g and the supernatant was filtered through Miracloth into another tube and centrifuged as before. The supernatant was again filtered through Miracloth into 15 mls of room temperature isopropanol in an Oak Ridge tube. After gentle mixing, the sample was incubated at room temperature for 10-60 minutes until the DNA 20 precipitated. The DNA was spooled and allowed to air dry before being resuspended in 4 mls of TE on ice for 1 hour. CsCl was added to 0.97g/ml final concentration and 300 ul 10mg/ml ethidium bromide was also added before filling VTI80 quick seal tubes. The sample was centrifuged overnight at 225,000 x g overnight. The 25 DNA was extracted with water saturated butanol and enough water was added to bring the volume to 4 mls before adding 2 volumes

EtOH. The DNA was spooled, air dried and resuspended in 200 ul sterile water.

Northern and Southern Analysis

5 For Northerns, 10ug of total RNA was isolated from various tissues, separated by electrophoresis in 1.2% agarose-formaldehyde gels and transferred onto Nytran Plus membranes (Schleicher and Schuell). Hybridization conditions consisted of a solution containing 50% formamide (v/v), 5xSSC, 0.1% SDS, 5mM EDTA, 10X  
10 Denhardts solution, 25mM sodium phosphate pH6.5 and 250 ug/ml carrier DNA. Washes were performed in 2xSSC, 0.1% SDS at 42°C 3 times for 30 minutes each time.

Cotton genomic DNA (12ug) was digested with various restriction endonucleases, electrophoresed in 0.9% agarose gels  
15 and blotted onto Nytran Plus membranes. Hybridization and filter washing conditions for both the 3' specific and full-length cDNA insert probes were as described for Northern analysis.

Probes derived from 3'-untranslated regions were synthesized via oligonucleotide primers from the Rac13 cDNA, corresponding to  
20 bases 600-619 and 843-864 (Figure 4). Each set of primers was used in a polymerase chain reaction to synthesize copies of 3'-specific DNA sequences. These sequences were used as templates in the generation of single-stranded, <sup>32</sup>P-labeled probes off the antisense strand in a polymerase chain reaction. The full-length  
25 cDNA inserts for Rac13 were used as templates for double stranded, random primed probes using the Prime-It kit (Stratagene).

## Example 2

Isolation of cDNA Clones from Cotton

CDNA to the 4-4 clone was isolated from the cotton fiber library described above, and shown to express in fiber but not 5 other tissues. This sequence was not related to any known protein. Only 400 kb of encoding sequence was present in this clone, so the library was rescreened using the cDNA to obtain full-length clones. The full-length encoding sequence is provided in Figure 1.

10 By comparing sequences of random cDNA clones against various sequence data banks via BLAST, a National Center for Biotechnology Information service, a clone, designated #105, was found to have an encoding sequence related to that of a reported lipid transfer protein.

15 Another clone was sequenced which showed high homology to animal Rac proteins. This clone, designated Rac, was not quite full-length, and the library was re-screened using this initial Rac DNA segment as probe. Of approximately 130,000 primary plaques screened, 56 screened positive; of these, 14 clones were 20 isolated and sequenced. Of these 14 clones, 12 showed identical sequence homology to the original Rac clone and one of these cDNA clones encoded a full length cDNA and received the name Rac13. Figure 4 shows the cDNA sequence encoding the Rac13 gene expressed in cotton fiber.

25 One other partial-length cDNA clone, designated Rac9, was clearly related, but distinct in DNA and amino acid sequence from Rac13. Re-screening of 150,000 plaques resulted in the isolation

of 36 positive clones of which only two clones corresponded to the Rac9 sequence (both full-length clones), the remainder being Rac13. These results suggest that cotton contains genes for at least two distinct Rac proteins. Based upon the frequency of 5 clone isolation, Rac13 is relatively highly-expressed and Rac9 less so in cotton fibers at 21 days post-anthesis (dpa), the age at which polyA<sup>+</sup> mRNA was isolated for library construction.

Comparisons of the deduced amino acid sequence of Rac13 with other small G-proteins showed that the cotton Rac proteins are 10 very closely related to the Rhol protein sequence deduced from a cDNA clone isolated recently from pea (Yang and Watson, *supra*). After the pea Rhol, mammalian Rac proteins show the highest homology with the cotton Rac proteins. Other proteins of the rho subfamily, such as the yeast CDC42 and human RhoA, are also 15 clearly related to the cotton Rac genes. By contrast, the other small G-proteins of the Rab/YPT subfamily isolated from plants such as the example shown of the tobacco RAB5 protein, as well as the human Ras proteins, are least homologous to the cotton Rac proteins of all the small G-proteins compared. The cotton and pea 20 proteins, as well as the mammalian Racs, all have pI's above 9, whereas those of other rho and ras proteins are in the range of 5.0-6.5.

### Example 3

#### 25 Expression of Cotton Fiber Genes in Developing Fibers

Expression of the Rac13 and 4-4 genes was assessed using mRNA prepared from various cotton tissues and from fibers at

different stages of development. Blots were hybridized with probes derived from untranslated regions of Ltp, Rac13 and 4-4 genes. The gene for Rac13 exhibits highly-enhanced expression in fibers; virtually no detectable mRNA is present in leaves, roots, 5 or flower parts, even under conditions of extended development time. Rac13 expression is detected in seeds at an age that corresponds to the highest expression levels observed in fiber tissue derived from seeds of this same age. The pattern of Rac13 expression in fibers is very dependent upon the developmental 10 stage. Expression is very low during the stage of primary wall synthesis (0-14 dpa, see Meinert and Delmer, 1977), reaches a maximum during the transition to secondary wall synthesis (about 15-18 dpa), and declining during the stage of maximal secondary wall cellulose synthesis (about 24-28 dpa).

15 4-4 mRNA begins to accumulate in fiber cells only at day 17 post anthesis and continues through at least day 35 post anthesis. Levels peak at day 21 and remain high. 4-4 mRNA is not detected in other cotton tissues, and is not detected in fiber tissue before onset at 17 days post anthesis.

20 The #105 lipid transfer protein cDNA clone was used as a probe against cotton tissue and in a cotton fiber northern. The northern showed that the cotton fiber Ltp is highly expressed in cotton fiber. The mRNA that codes for this protein is expressed throughout fiber development at extremely high level. Northern 25 blots indicate that this mRNA is expressed at 5 dpa and is continually expressed at a high level at 40 dpa.

## Example 4

Genomic DNA

cDNA for both the 4-4 and Rac13 was used to probe for genomic clones. For both, full length genomic DNA was obtained 5 from a library made using the lambda dash 2 vector from Stratagene™, which was used to construct a genomic DNA library from cotton variety Coker 130 (*Gossypium hirsutum* cv. coker 130), using DNA obtained from germinating seedlings.

The cotton genomic library was probed with a 3'-specific Ltp probe and 6 genomic phage candidates were identified and purified. 10 Figure 7 provides an approximately 2 kb sequence of the Ltp promoter region which is immediately 5' to the Ltp encoding region.

Six genomic phage clones from the cotton genomic library 15 were identified using a 3'-specific probe for the Ltp mRNA. This was done to select the promoter from the Ltp gene that is maximally expressed in cotton fiber from the family of Ltp genes in cotton. The Ltp promoter is active throughout the fiber development period.

20

## Example 5

Preparation of 4-4 Promoter ConstructspCGN5606

The pCGN5606 promoter construct comprises the 4-4 cotton 25 fiber expression cassette in a first version, version I (Figure 2). The sequences from nt1 to 65 and nt 5,494 to 5,547 correspond to fragments of the pBluescriptII polylinker where this cassette

is cloned. Unique restriction enzyme sites present in these regions flanking the cassette allow the cloning of the fiber expression cassette into binary vectors including the pCGN 5138 and 1547 series.

5 The sequences from nt57 to 5,494 are contained in a lambda phage clone of a cotton Coker 130 genomic library. This lambda genomic clone was given the designation 4-4(6).

The region from nt 65 to nt 4,163 corresponds to the 5' flanking region of the 4-4(6) gene. At nt 4,163 there is a NcoI 10 restriction site sequence that corresponds to the first codon of the 4-4 (6)ORF.

The region from nucleotide 4,163 to 4,502 corresponds to part 15 of the 4-4 (6)ORF. The sequence from nt 4,502 to 4,555 is a synthetic polylinker oligonucleotide that contains unique target sites for the restriction enzymes EcoRI, SmaI, SalI, NheI and BglII. This fragment from nt4,163 to 4,555 is a stuffer fragment and is left in place to facilitate the monitoring of cloning 20 manipulations.

The genes to be expressed in cotton fiber cells using this cassette can be cloned between the NcoI restriction site and any 25 of the polylinker sites. This operation will replace the stuffer fragment with the gene of interest. The region from nt 4,555to 5,494 corresponds to the 940 nucleotides downstream of the stop codon and constitute the 3' flanking region of the 4-4 (6) gene. There is a unique AscI restriction enzyme site at nt 5483.

pCGN5610

The pCGN5610 construct is a second version of a 4-4 cotton fiber expression cassette, version II, which is a modified version of pCGN5606. The two versions of the 4-4 cotton fiber expression cassette are designed to allow the cloning of tandem arrays of two fiber cassettes in one binary plasmid. The differences with respect to pCGN5606 are very minor and described below.

The XbaI restriction site in the region of nt 1 to 65 has been deleted by standard cloning manipulations.

The polylinker region is in the reverse orientation of pCGN5606.

10 There is a unique XbaI restriction enzyme site at nt5484. The sequences from nt1 to 57 and nt 5,494 to 5,518 of pCGN5610 correspond to fragments of the pBluescriptII polylinker where this cassette is cloned. Unique restriction enzyme sites present in these regions allow the cloning of the fiber expression cassette  
15 into binary vectors of the pCGN 5138 and 1547 series.

The sequences from nt57 to 5,494 are contained a lambda phage clone of a Coker 130 genomic library. This clone is described in my notebook as lambda genomic clone 4-4(6). The region from nt 57 to nt 4,155 corresponds to the 5' flanking 20 region. At nt 4,155 there is a NcoI restriction site sequence that corresponds to the first codon of the 4-4 ORF. The region from nucleotide 4,156 to 4,500 corresponds to part of the 4-4 ORF. This fragment from nt4,156 to 4,550 is a stuffer fragment and is left in place to facilitate the monitoring of 25 cloning manipulations. The sequence from nt 4,500 to 4,550 is a synthetic polylinker oligonucleotide containing unique target

sites for the restriction enzymes BglII, NheI, SalI, SmaI and EcoRI.

The genes to be expressed in cotton fiber cells using this cassette can be cloned between the NcoI restriction site and any 5 of the polylinker sites. This operation replaces the stuffer fragment with the gene of interest. The region from nt 4,550 to 5,494 corresponds to the 940 nucleotides downstream of the stop codon and constitute the 3' flanking region of the 4-4 (6) gene.

10

#### Example 6

##### Preparation of Rac13 Promoter Constructs

###### Genomic clone

From a genomic clone designated 15-1, mapping was done with restriction endonucleases. The largest fragment with the Rac13 15 coding region was identified. This was a Pst fragment, and when subcloned in the Bluescript™ KS+ vector (BSKS+; Stratagene) was named pCGN4722. The insert had a length of 9.2 kb.

The region of the Pst fragment with the Rac13 coding sequence was identified. DNA sequence was determined for approximately 1.7 20 kb 5' of the start codon and approximately 1.2 kb 3' of the stop codon. The entire Rac coding region (exons and introns) was conveniently flanked by NdeI sites.

pCGN4722 was digested with XbaI, and a 2.7 kb fragment was removed. Religation gave pCGN4730, which was then digested with 25 NdeI, dropping out a 1.7 kb fragment containing the entire Rac coding region. Religation yielded pCGN4731.

A polylinker region was created using overlapping synthetic oligonucleotides which were PCR'ed using primers homologous to the 5' and 3' ends of the resynthesized section. The resulting product was digested with EcoRI and Hind III and ligated into 5 BSKS+ at the EcoRI and Hind III sites. The resulting plasmid was designated pCGN4733.

pCGN4731 and pCGN4633 were digested with NdeI and the NdeI fragment containing the synthesized polylinker region from pCGN4733 was dropped in the NdeI site of 4731, giving pCGN4734. 10 This last plasmid was digested with Sal and Xba, and so was pCGN5133. pCGN5133 was the 9.2 kb pst fragment in BSKS+ where the polylinker sites flanking the insert were altered to different sites for ease of manipulation. The fragment from pCGN4734 was then placed into the equivalent site of pCGN5143, giving pCGN4735.

15 A sequence for approximately 3 kb of the promoter construct pCGN4735 is provided in Figure 5. The resynthesized sequence falls between the NdeI sites located at bases 1706 and 1898 of the sequences. Thus, the sequence in Figure 5 includes approximately 1.7 kb 5' to the NdeI site 5' to the resynthesized polylinker 20 region. There is a roughly 2.5 kb sequence 5' from this sequence which is not provided in Figure 5, relative to the total 9.2 kb insert. The sequence of Figure 5 also includes approximately 1.1 kb 3' to the 3' NdeI site. Approximately 3 kb which is most 3' in the Rac13 insert is not provided in Figure 5. A map for pCGN4735 25 is provided in Figure 6.

Example 7

Pigment Synthesis GenesMelanin

A binary construct for plant transformation to express genes for melanin synthesis is prepared as follows. The melanin genes were originally isolated from the common soil bacterium *Streptomyces antibioticus* (Bernan et al. (1985) 34:101-110). Melanin production is composed of a two gene system. The first gene, *tyrA*, encodes the catalytic unit responsible for the polymerization of the amino acid tyrosine, the primary substrate, and is termed tyrosinase. The second gene, ORF438, is responsible for binding copper and delivering copper to the tyrosinase and activating the enzyme. Expression of both the ORF438 and *tyrA* genes ensures maximal tyrosinase activity.

The genes for both ORF438 and *tyrA* were fully re-synthesized with respect to their DNA sequence. This was performed as the initial DNA sequence isolated from *Streptomyces* has a very high guanine and cytosine (G+C) DNA content. Thus, the ORF438 and *tryA* genes were re-synthesized to appear more "plant-like" (reduced G+C content) with respect to plant preferred codons encoding their corresponding amino acids.

Indigo

Indigo production involves conversion of the amino acid tryptophan, the primary substrate, into indole which is then converted into indoxyl. Molecules of indoxyl spontaneously convert to indigo in the presence of oxygen. A two gene system was used to affect indigo production in fiber cells. The first

gene (*tna*) was obtained from the bacterium *E. coli* and encodes the enzyme tryptophanase. The designation *tta* stands for the gene encoding tryptophanase from *E. coli*, an enzyme which converts tryptophan to indole (Stewart et al., (1986) *J Bacteriol* 166:217-5 223).

The *pig* designation is used for the encoding sequence to the protein for indigo production from *Rhodococcus*, which produces indigo from indole (Hart et al., (1990) *J Gen Microbiol* 136:1357-10 1363). Both *tta* and *pig* were obtained by PCR. Tryptophanase is responsible for the conversion of tryptophan to indole, while the second gene (*pig*) encodes an indole oxygenase enzyme responsible for the conversion of indole to indoxyll. Both these bacterial genes were utilized in their native form.

15

Example 8Constructs for Targeting Pigment Synthesis Genes

For plastid targeting, the constructs contain a fragment of the tobacco ribulose bisphosphate carboxylase small subunit gene encoding the transit peptide and 12 amino acids of the mature 20 protein (Tssu) positioned in reading frame with the appropriate encoding sequence.

For vacuolar targeting of the melanin synthesis genes, constructs include a fragment of the metallocarboxypeptidase inhibitor gene, encoding the entire 32 amino acid N-terminus 25 signal peptide of that protein plus 6 amino acids of the mature protein (CPI+6) (Martineau et al., *supra*), positioned in reading frame with the appropriate encoding sequences. In addition to the

signal peptide, a sequence encoding a vacuolar localization signal (VLS) is inserted 3' of the protein encoding sequence.

Constructs which contain encoding sequences for bacterial genes involved in biosynthesis of pigmented compounds and 5 sequences for directing transport of the encoded proteins into plastids or vacuoles are prepared as follows.

Melanin

The re-synthesized ORF438 and *tyrA* genes were treated in two 10 distinct ways depending on which compartment in the fiber cell the final protein products would be localized. One chimeric gene/plant binary construct (designated pCGN5148) contained the genes targeted to the fiber cell plastids. To do this, 12 amino acids of a gene for the small subunit of carboxylase (SSU) plus 15 the original 54 amino acid SSU transit peptide were fused to the amino termini of both the ORF438 and *tyrA* gene products respectively. These peptide sequences allow the ORF438 and *tyrA* gene products (proteins) to be efficiently targeted to the plastid. This targeting was initiated as the plastid is the site 20 of tyrosine production within the fiber cell.

The second chimeric gene/plant binary construct (designated pCGN5149) contained the ORF438 and *tyrA* genes targeted to the vacuole within the fiber cell. Based on information from other biological systems, it was postulated that the fiber cell vacuole 25 may contain a high concentration of tyrosine for melanin polymerization. Both the ORF438 and *tryA* genes contain the 29 amino acid signal peptide from a tomato carboxypeptidase inhibitor

(CPI) protein as amino terminal gene fusions to direct these proteins to the endoplasmic reticulum (ER) secretory system of the fiber cell.

In addition, the *tyrA* gene has an 8 amino acid vacuolar targeting peptide (VTP) from CPI fused at the carboxy terminus so that the mature copper-activated tyrosinase will eventually be targeted to the vacuole of the fiber cell. Both the ORF438 and *tyrA* proteins also had potential glycosylation sites removed via site-directed mutagenesis of the ORF438 and *tyrA* genes respectively. Potential plant cell glycosylation of these proteins upon their expression in fiber cells could result in tyrosinase inactivation, hence removal of potential glycosylation sites was deemed necessary.

15 Indigo

The only modification to the indigo genes was the fusion of the tobacco SSU transit peptide encoding DNA sequences onto the amino terminal region of both the *tua* and *pig* genes to affect the localization of both the tryptophanase and indole oxygenase proteins to the fiber cell plastid. These are the same exact gene fusions that were made for the plastid-directed proteins for melanin production in construct 5148. The *tua* and *pig* gene products were targeted to the fiber cell plastid as that is the primary site of tryptophan synthesis.

25

Example 9

Expression Constructs

Melanin

The modified genes for both the plastid and vacuolar targeted ORF438 and tyrosinase proteins were placed into a fiber expression cassette to be "switched" on during development of the cotton 5 fiber cell. The "switch" (promoter) utilized for the melanin constructs was 4-4. The modified ORF438 and *tyrA* genes were cloned into the 4-4 promoter cassette and these chimeric genes then inserted into a binary plasmid to create plasmids pCGN5148 and pCGN5149, containing the modified genes for plastid and 10 vacuolar targeted ORF438 and tyrosinase proteins, respectively. These binary plasmids also contain genetic determinants for their stable maintenance in *E. coli* and *Agrobacterium* and also contain a chimeric gene for plant cell expression of the bacterial kanamycin 15 resistance gene. This kanamycin resistance marker allows for the selection of transformed versus non-transformed cotton cells when plant hypocotyl or leaf segments are infected with *Agrobacterium* containing the binary plasmids.

A block diagram of the plasmid pCGN5149, having vacuolar targetting sequences, is shown in Figure 8. Plasmid pCGN5148 (not 20 shown) is constructed the same as 5149, only pCGN5148 has plastid-targetting sequences.

Indigo

As with the melanin genes, the plastid-directed *tta* and *pig* 25 genes were placed in the fiber-specific 4-4 promoter cassette and these chimeric genes subsequently inserted into a binary plasmid

to create plasmid pCGN5616. A block diagram of plasmid pCGN5616 is shown in Figure 8.

#### Anthocyanin

5       A construct has been prepared for the expression of the maize R and CI genes in developing cotton fiber. These genes are known to be responsible for the production of Anthocyanin pigments by acting in a regulatory manner to turn on the chalcone pathway for production of anthocyanins (red spectrum colors). The R and CI 10 genes were placed under the control of the Rac13 promoter cassette. A binary plasmid designated pCGN4745 (not shown), contains both the R and CI genes each under control of the Rac13 promoter.

15

#### Example 10

##### Cotton Transformation

###### Explant Preparation

20       Coker 315 seeds are surface disinfected by placing in 50% Clorox (2.5% sodium hypochlorite solution) for 20 minutes and rinsing 3 times in sterile distilled water. Following surface sterilization, seeds are germinated in 25 x 150 sterile tubes containing 25 mls 1/2 x MS salts: 1/2 x B5 vitamins: 1.5% glucose: 0.3% gelrite. Seedlings are germinated in the dark at 28°C for 7 days. On the seventh day seedlings are placed in the light at 25 28±2°C.

###### Cocultivation and Plant Regeneration

Single colonies of *A. tumefaciens* strain 2760 containing binary plasmids pCGN2917 and pCGN2926 are transferred to 5 ml of MG/L broth and grown overnight at 30°C. Bacteria cultures are diluted to 1 x 10<sup>8</sup> cells/ml with MG/L just prior to cocultivation.

- 5 Hypocotyls are excised from eight day old seedlings, cut into 0.5-0.7 cm sections and placed onto tobacco feeder plates (Horsch et al. 1985). Feeder plates are prepared one day before use by plating 1.0 ml tobacco suspension culture onto a petri plate containing Callus Initiation Medium CIM without antibiotics (MS salts: B5 vitamins: 3% glucose: 0.1 mg/L 2,4-D: 0.1 mg/L kinetin: 0.3% gelrite, pH adjusted to 5.8 prior to autoclaving). A sterile filter paper disc (Whatman #1) was placed on top of the feeder cells prior to use. After all sections are prepared, each section was dipped into an *A. tumefaciens* culture, blotted on sterile paper towels and returned to the tobacco feeder plates.
- 15

- Following two days of cocultivation on the feeder plates, hypocotyl sections are placed on fresh Callus Initiation Medium containing 75 mg/L kanamycin and 500 mg/L carbenicillin. Tissue was incubated at 28±2°C, 30uE 16:8 light:dark period for 4 weeks.
- 20 At four weeks the entire explant was transferred to fresh callus initiation medium containing antibiotics. After two weeks on the second pass, the callus was removed from the explants and split between Callus Initiation Medium and Regeneration Medium (MS salts: 40mM KNO<sub>3</sub>: 10 mM NH<sub>4</sub>Cl:B5 vitamins:3% glucose:0.3% gelrite:400 mg/L carb:75 mg/L kanamycin).
  - 25

Embryogenic callus was identified 2-6 months following initiation and was subcultured onto fresh regeneration medium.

Embryos are selected for germination, placed in static liquid Embryo Pulsing Medium (Stewart and Hsu medium: 0.01 mg/l NAA: 0.01 mg/L kinetin: 0.2 mg/L GA3) and incubated overnight at 30°C. The embryos are blotted on paper towels and placed into Magenta boxes 5 containing 40 mls of Stewart and Hsu medium solidified with Gelrite. Germinating embryos are maintained at  $28\pm 2^\circ\text{C}$   $50 \mu\text{E m}^{-2}\text{s}^{-1}$  16:8 photoperiod. Rooted plantlets are transferred to soil and established in the greenhouse.

Cotton growth conditions in growth chambers are as follows: 10 16 hour photoperiod, temperature of approximately  $80-85^\circ$ , light intensity of approximately  $500\mu\text{Einstins}$ . Cotton growth conditions in greenhouses are as follows: 14-16 hour photoperiod with light intensity of at least  $400\mu\text{Einstins}$ , day temperature  $90-95^\circ\text{F}$ , night temperature  $70-75^\circ\text{F}$ , relative humidity to 15 approximately 80%.

#### Plant Analysis

Flowers from greenhouse grown T1 plants are tagged at anthesis in the greenhouse. Squares (cotton flower buds), 20 flowers, bolls etc. are harvested from these plants at various stages of development and assayed for enzyme activity. GUS fluorometric and histochemical assays are performed on hand cut sections as described in co-pending application filed for Martineau et al., *supra*. For fiber color characteristics, plants 25 are visually inspected, or northern or western analysis can be performed, if necessary.

Example 11Expression of Transgenic Pigment Synthesis GenesMelanin

5 Plants that exhibited resistance to the kanamycin selectable marker via a leaf assay and corresponding Western analysis were considered transformed. Transgenic fiber was collected from individual plant transformants at different stages of fiber development and analyze in two ways. One was to analyze fiber at  
10 a single developmental time point for each transgenic cotton plant to compare tyrosinase expression between transgenic events. The other was to screen developing fiber from selected plants to analyze the timing of tyrosinase expression under the control of the fiber-specific 4-4 promoter, by Western blots using antisera  
15 prepared against purified tyrosinase protein.

For the plastid-targeted construct pCGN5148 9 of 13 events screened for tyrosinase expression were positive, while 13 of the 16 transformed vacuolar-targeted construct pCGN5149 events which were screened were positive. Expression level in the fiber in  
20 tyrosinase positive plants is approximately 0.1-0.5% fiber cell protein. Clearly, the cotton fiber cells comprising the DNA color constructs DNA produce the necessary proteins required for synthesis of a pigment.

Visually, the lint from the tyrosinase positive events  
25 exhibits color to varying degrees, while plants that do not express the enzyme do not exhibit any color. Colorimeter measurements of cotton fiber taken from control Coker 130 plants

and plants from various events transformed with pCGN5148 are provided in Figures 9 and 10, respectively.

Fiber from pCGN5148 (plastid-directed) plants demonstrates a bluish-green color phenotype. One event, 5148-50-2-1 included 5 cotton fiber cells (linters) which were colored and which had an negative  $a^*$  value less than - 8.0, as measured on the  $L^*a^*b^*$  color space. Coker 130 cotton fiber cells do not typically demonstrate a negative  $a^*$  value.

These colored cotton cells also had a color located on the 10  $L^*C^*h$  color space with a relatively high hue angle value  $h$ , greater than  $135^\circ$ . Normal Coker 130 fibers have a similar value which is not greater than about  $90^\circ$  as measured by this method.

Results of colorimeter measurements of cotton fiber taken from plants transformed with pCGN5149 are provided in Figure 11. 15 Fiber from plants expressing tyrosinase from construct pCGN5149 (vacuolar-targetted) tends to have a light brown phenotype.

#### Indigo

Resistance to the kanamycin selectable marker via leaf assay 20 and Western analysis was again the criterion for designating a plant as transformed by pCGN5616. Transgenic fiber was collected from individual plant transformants at different stages of fiber development. The transgenic developing fiber is screened from selected plants to analyze the timing of *tua* and *pig* gene 25 expression under the control of the fiber-specific 4-4 promoter and fiber is also analyzed at a single developmental time point for each transgenic cotton plant for comparison of both

tryptophanase and indole oxygenase expression between transgenic events, by using Western blots with antisera prepared against the tryptophanase and indole oxygenase proteins.

For the indigo events, 15 of 24 screened plants were positive for expression of both the tryptophanase and indole oxygenase enzymes. Expression levels in the fiber of these proteins is between 0.05-0.5% fiber cell protein. Approximately half of these transformants are expressing both genes in the fiber resulting in a very faint light blue color phenotype. Visually, there is a faint blue color in the majority of these positive events, particularly in 20-30 dpa fiber in the unopened boll. Results of colorimeter measurements of cotton fiber taken from various events of plants transformed with pCGN5616 are provided in Figure 12. Many of these events had relatively low  $a^*$  values (less than 2) with elevated  $b^*$  values (greater than 10), as measured on the L\* $a^*b^*$  color space. Similarly, several 5149 events also measured with an  $a^*$  value less than 2 while maintaining a  $b^*$  value greater than 10.

20 BC Cotton

Colorimeter measurements taken on naturally colored fiber from four separate BC cotton lines is provided in Figure 13.

The above results demonstrate that the color phenotype of a transgenic cotton fiber cell can be altered by expressing pigment synthesis genes. The transgenic cotton fiber cells include both a pigment synthesizing protein, and pigment produced by the pigment

synthesizing protein. As shown from the results of Figures 9 through 13, expression of a pigment gene of interest can result in cotton fiber cells in which the synthesis of pigments combined with appropriate targeting sequences results in modification of 5 color phenotype in the selected plant tissue, yielding colored cotton fiber by expression from a genetically engineered construct.

All publications and patent applications cited in this 10 specification are herein incorporated by reference as if each individual publication or patent application are specifically and individually indicated to be incorporated by reference.

Although the foregoing invention has been described in some detail, by way of illustration and example for purposes of clarity 15 and understanding, it will be readily apparent to those of ordinary skill in the art that certain changes and modifications may be made thereto, without departing from the spirit or scope of the appended claims.

CLAIMS

What is claimed is:

1. A DNA construct comprising as operably joined  
5 components in the direction of transcription, a cotton fiber  
transcriptional factor and an open reading frame encoding a  
protein of interest, wherein said transcriptional factor is  
selected from the group consisting of the Ltp, the 4-4 and  
the rac promoter sequences.

10 2. The DNA construct according to Claim 1, further  
comprising a transport signal encoding sequence from a plant  
nuclear-encoded gene.

15 3. The DNA construct according to Claim 2, wherein said  
transport signal encoding sequence comprises a plastid  
transit peptid.

4. The DNA construct according to Claim 1, wherein said  
transport signal encoding sequence encodes a signal peptide  
which provides for transport across the rough endoplasmic  
reticulum.

20 5. The DNA construct according to Claim 4, wherein said  
sequence further comprises, 3' to said open reading frame, a  
vacuolar localization signal.

6. The DNA construct of Claim 1 wherein said pigment is  
melanin or indigo.

25 7. The DNA construct of Claim 6 wherein said open  
reading frame is from a bacterial gene.

8. The DNA construct of Claim 7 wherein said bacterial gene is selected from the group consisting of ORF438, *tyrA*, anthocyanin R gene, anthocyanin C1 gene, *pig*, and *tta*.

9. A plant cell comprising a DNA construct of Claim 1.

5 10. A cotton plant cell according to Claim 9.

11. A cotton fiber cell according to Claim 10.

12. A plant comprising a cell of any one of Claims 9-  
11.

13. A method of modifying fiber phenotype in a cotton  
10 plant, said method comprising:

transforming a plant cell with DNA comprising a construct for expression of a protein in a pigment biosynthesis pathway, wherein said construct comprises as operably joined components:

15 a transcriptional initiation region functional in cells of said cotton plant,

an open reading frame encoding a protein of interest,  
and

20 a transcriptional termination region functional in cells of said cotton plant,

wherein said plant cell comprises a substrate of said protein; and

growing said plant cell to produce a cotton plant,  
wherein said protein reacts with said substrate to produce  
25 said pigment.

14. The method of Claim 13 wherein said construct further comprises a transport signal encoding sequence from a plant nuclear-encoded gene.

15. The method of Claim 13 wherein said transport signal encoding sequence encodes a signal peptide which provides for transport across the rough endoplasmic reticulum.

16. The method of Claim 13 wherein said DNA comprises constructs for expression of two proteins in a pigment biosynthesis pathway, wherein each of said constructs comprises components i) through iv), and wherein said two proteins are not encoded by the same gene.

17. The method of Claim 16 wherein said pigment is melanin and said proteins are encoded by *tyrA* and *ORF438*.

18. The method of Claim 16 wherein said pigment is indigo and said proteins are *tna* and *pig*.

19. The method of Claim 16 wherein said pigment is anthocyanin and said constructs comprise the anthocyanin R and C1 regulatory genes.

20. The method of Claim 13 wherein plant cell is a cotton fiber cell, and wherein said transcriptional region is a fiber tissue transcription initiation region.

21. The method of Claim 20 wherein said transcriptional region is selected from the group consisting of the *Ltp*, the 4-4 and the *rac* promoter sequences

22. A recombinant DNA construct comprising the cotton tissue transcriptional sequence shown in Figure 2.

23. A recombinant DNA construct comprising the cotton tissue transcriptional sequence shown in Figure 5.

24. An isolated DNA encoding sequence of Figure 1.

25. An isolated DNA encoding sequence of Figure 4.

5 26. The method of Claim 13 wherein said protein of interest is involved in the synthesis of a plant hormone.

27. An isolated DNA sequence comprising the cotton lipid transfer protein encoding sequence of Figure 7.

10 28. A cotton fiber cell comprising a DNA sequence, wherein said DNA sequence comprises as operably joined components in the direction of transcription, a cotton fiber transcriptional factor and an open reading frame encoding a protein required for synthesis of a pigment.

15 29. A cotton fiber cell according to Claim 27 comprising pigment produced by said pigment synthesizing protein.

30. A cotton fiber cell according to Claim 27 wherein said DNA sequence further comprises a transport signal encoding a sequence from a plant nuclear-encoded gene.

20 31. A cotton fiber cell according to Claim 29, wherein said transport signal encoding sequence comprises a plastid transit peptid.

32. A cotton fiber cell according to Claim 29, wherein said transport signal encoding sequence encodes a signal peptide which provides for transport across the rough endoplasmic reticulum.

25 33. A cotton fiber cell according to Claim 31, wherein said sequence further comprises, 3' to said open reading frame, a vacuolar localization signal.

34. A cotton fiber cell according to Claim 27 wherein said transcriptional factor is selected from the group consisting of the cotton fiber lipid transfer promoter sequence, the 4-4 promoter sequence and the rac promoter sequence.

5 35. A cotton fiber cell according to Claim 27 wherein said pigment is melanin or indigo.

36. A cotton fiber cell according to Claim 27 wherein said open reading frame is from a bacterial gene.

10 37. A cotton fiber cell according to Claim 35 wherein said bacterial gene is selected from the group consisting of ORF438, tyrA, anthocyanin R gene, anthocyanin C1 gene, pig, and tna.

38. A cotton fiber cell comprising melanin.

39. A cotton fiber cell comprising indigo.

15 40. A cotton fiber cell which is colored by genetic engineering and which has a negative a\* value less than - 1.0 as measured on the L\*a\*b\* color space.

41. The cotton fiber cell of Claim 39 wherein said negative a\* value is less than a -5.0.

20 42. The cotton fiber cell of Claim 40 wherein said negative a\* value is less than a -8.0.

43. A cotton fiber cell which is colored by genetic engineering and which has an a\* value less than 2 and the b\* value greater than 10 as measured on the L\*a\*b\* color space.

25 44. A cotton fiber cell which is colored by genetic engineering and which has a hue angle value h of greater than 100° as measured on the L\*C\*h color space.

45. The cotton fiber cell of Claim 43 wherein said h value is greater than a 135°.

CTT TCT ATT TGG TTA ACC ATG GCT CAT AAC TTT CGT CAT CCT TTC  
 Leu Ser Ile Trp Leu Thr Met Ala His Asn Phe Arg His Pro Phe Phe>  
 20  
 40

CTT TTC CAA CTT TTA CTC ATT ACT GTC TCA ATG ATC GGT AGC CAC  
 Leu Phe Gln Leu Leu Ile Thr Val Ser Leu Met Ile Gly Ser His>  
 60  
 80

ACC GTC TCG TCA GCG GCT CGA CAT TTA TTC CAC ACA CAA ACA ACC TCA  
 Thr Val Ser Ala Ala Arg His Leu Phe His Thr Gln Thr Thr Ser>  
 100  
 120  
 140

TCA GAG CTG CCA CAA TTG GCT TCA AAA TAC GAA AAG CAC GAA GAG TCT  
 Ser Glu Leu Pro Gln Leu Ala Ser Lys Tyr Glu Lys His Glu Glu Ser>  
 160  
 180

GAA TAC AAA CAG CCA AAA TAT CAT GAA GAG TAC CCA AAA CAT GAG AAG  
 Glu Tyr Lys Gln Pro Lys Tyr His Glu Glu Tyr Pro Lys His Glu Lys>  
 200  
 220  
 240

CCT GAA ATG TAC AAG GAG GAA AAA CAA AAA CCC TGC AAA CAT CAT GAA  
 Pro Glu Met Tyr Lys Glu Glu Lys Gln Lys Pro Cys Lys His His Glu>  
 260  
 280

GAG TAC CAC GAG TCA CGC GAA TCG AAG GAG CAC GAA GAG TAC GAT AAA  
 Glu Tyr His Glu Ser Arg Glu Ser Lys Glu His Glu Glu Tyr Asp Lys>  
 300  
 320

GAA AAA CCC GAT TTC CCC AAA TGG GAA AAG CCT AAA GAG CAC GAG AAA  
 Glu Lys Pro Asp Phe Pro Lys Trp Glu Lys Pro Lys His Glu His Glu Lys>  
 340  
 360  
 380

400  
 420

\*

**FIGURE 1A**

CAC GAA GTC GAA TAT CCG AAA ATA CCC GAG TAC AAG GAC AAA CAA GAT His Glu Val Glu Tyr Pro Lys Ile Pro Glu Tyr Lys Asp Lys Glu Ser>	440	460	480
GAG AAT AAG AAA CAT AAA GAT GAA GAG TGC CAG GAG TCA CAC GAA TCG Glu Asn Lys Lys His Lys Asp Glu Glu Cys Gln Glu Ser His Glu Ser>	500	520	
AAA GAG CAC GAA GAG TAC GAG AAA GAA AAA CCC GAT TTC CCC AAA TGG Lys Glu His Glu Glu Tyr Glu Lys Glu Lys Pro Asp Phe Pro Lys Trp>	540	560	
GAA AAG CCT AAA GGG CAC GAG AAA CAT AAA GCC GAA TAT CCG AAA ATA Glu Lys Pro Lys Gly His Glu Lys His Lys Ala Glu Tyr Pro Lys Ile>	580	600	620
CCT GAG TGC AAG GAA AAA CTA GAT GAG GAT AAG GAA CAT AAA CAT GAG Pro Glu Cys Lys Glu Lys Leu Asp Glu Asp Lys Glu His Lys His Glu>	640	660	
TTC CCA AAG CAT GAA AAA GAA GAG GAG AAG AAA CCT GAG AAA GGC ATA Phe Pro Lys His Glu Lys Glu Glu Glu Lys Lys Pro Glu Lys Gly Ile>	680	700	720
GTA CCC TGA GTG GGT TAA AAT GCC TGA ATG GCC GAA GTC CAT GTT TAC Val Pro *** Val Gly *** Asn Ala *** Met Ala Glu Val His Val Tyr>	740	760	
TCA GTC TGG CTC GAG CAC TAA GCC TTA AGC CAT ATG ACA CTG GTG CAT Ser Val Trp Leu Glu His *** Ala Leu Ser His Met Thr Leu Val His>	780	800	*

FIGURE 1B

GTG CCA TCA TCA TGC AGT AAT TTC ATG GGA TAT TGT AAT TAT ATT GTT  
Val Pro Ser Ser Cys Ser Asn Phe Met Gly Tyr Cys Asn Tyr Ile Val>  
820  
AAT AAA AAA GAT GGT GAG TGG GAA ATG TGT GTG TGC ATT CAT CCA TGA  
Asn Lys Lys Asp Gly Glu Trp Glu Met Cys Val Cys Ile His Pro \*\*\*>  
840  
880  
GCA ATG CTG AAT CTC TTT GCA TGC ATA GAG ATT CTG AAT GGT TAT AGT  
Ala Met Leu Asn Leu Phe Ala Cys Ile Glu Ile Leu Asn Gly Tyr Ser>  
900  
920  
TTA TGT TAT ATC GTT TGT TCT AGT GAA ATT AAT TTT GAA TGT TGT ATG  
Leu Cys Tyr Ile Val Cys Ser Ser Glu Ile Asn Phe Glu Cys Cys Met>  
940  
TAA TGT T  
\*\*\* Cys Xxx>

FIGURE 1C

20 ACTAAAGGGA ACAAAAGCTG GAGCTCCACC CGGGTGGGG CCGCTCTAGA ACTAGTGGAT  
 80 \* 100 120  
 CCCCGGTGGA CTAACAAAAA CATGGAAAGA TTGCTGTTAA AAAATAAAA GAAGCTTACT  
 140 160 180  
 CAATAACACT TTGTAATG TATACAAAG ACTCAATGAA AAACAATAAC TCAATAACACT  
 200 220 240  
 TTTTTCAC TATTACATC CTTTATATAG GCTGAAACTA CAACAACTT AGCTAAAAAA  
 260 280 300 \*  
 ATAGGATAAC CTAATAGCAA AATCACAATC AGATATTTAA CCATGATT AGCTAACCAT  
 320 340 360  
 TTAACAACTT TATTGAAACT AATTGAAATA TTTCATCTGC TGATATGCC AAGATTAG  
 380 400 420  
 GCCACTAACC GATTGGTGG TGAACTTTAA CATGTCATGC ATTGTAAC GTTGAAACA  
 440 460 480  
 AGTTTTTGC ATTATTTAC TATATGAACT GTTGTGTTAG GTTGAGTAC ACACTGAGCT  
 500 520 540  
 TGTAAAGCTCA CTCAAATT TTCTAATTCTT AAGGTGATCA GCAAACCTAG GACGGGGGG  
 560 580 600 \*  
 CGTACGGAGAG CTCGGATTGA TTCTCTAGTT AATAATAAG ACGATTATG TTTTAACT

Figure 2A

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ATTATGGACT TTTGGACTA TGTAAC TGTGTT TGGGACTTAA TTTTTGTTT TTATTGCTT  
 620 640 660  
 680 700 \* 720  
  
 TTTTGAGATT TAGTAATTAT TATTTTAAA CTGCCAAATT ATATGTTTT ACAAACTAAG  
  
 TCACAGTTT CAAAATTCCA TAACCTGAA TTTTTCGCTG CAAAATAAG TAATCATTAA  
 740 760 780  
  
 800 820 840  
  
 \*  
  
 AGTGTTTTT CTGTAATAAA ATAATAAAAT AATTAAACG AGTATTTCG TAAATAATTGG  
  
 860 880 900 \*  
  
 AAATTGATT ACCAAATAA GTATGTCAAA ACACATGTT ATATGTTACA GGGCGATATC  
  
 GTCTAGGCAA ATAACATCTA GGCGGGTTT GGAGTGTAC AGGGCGAGTG GGCTCATTT  
  
 920 940 960  
  
 980 1000 \* 1020  
  
 GAGTAAGTAT AGTTAGGGCC GAGTTTAGA TTGCATATTCA AAGGTCAAAG ATTTGTAAA  
  
 CTTCGATGAA TGATATGTAT GATTGTCGGA TAAACGAAAT ATGTTTTTTT CTTTGTGTC  
  
 1040 1060 1080  
  
 1100 1120 1140  
  
 \*  
  
 TGTTTATCT CGTGTGATAA GTATATAGTA TGTTTATTCA CAATTCTTAT GGCATGTGAC  
  
 1160 1180 1200 \*  
  
 ATTGTGGCTA TTCTAATTAA ATTGATTGTT TATTATGAA ATCTGATGCA TCTGTTCTAC  
  
 1220 1240 1260

Figure 2B

AAAGCATGGA ATCTCATGCC	TACTGCCTTC	TGTTAAAGAT ACGATTGCAA	GTTAACATG
1280	1300	*	1320
CTTACTATTT TGATTTGTC	CTTGCATGCT	ATGTACACATT ACATGGGGTT	GGCATGATAT
1340	1360		1380
GGTAAGGAGG AAGTTTTGAC	AGTTAAATGA	TTTGCACTAT CTGGTGGTT AACACATAT	
1400	1420		1440
TTGTTATGGC ATCTTGACTG	CGGTTATGGT	GGCTCGACCG CCCATATCTG	TTCTGGAAAT
1460	1480	*	1500
TTATCTGTGA CTCTGGGGC	ATTGTCTACA	ATTATTGTT GGATGGACGA	
1520	1540		1560
GTCGTGGGA ACTCTATTG	GTGTGTTGCG	GAGTTGGTA GGAATTTTC	AAAAAAATT
1580	1600	*	1620
TGCATTGCT GTTTCTGAAA	AATATTGCAT	TAACATAATC ATGCATTCTC	AATTGGTC
1640	1660		1680
AATTGAAACGT TATAAAATTC	TCTATGATAT	CCTGATCTGT TTATTACATT	ATATGTGTT
1700	1720	*	1740
ATGCTTGACT TAAGTCAAAC	ATTGAGATTG	ATAGCTCACC	CAATTATTA ATCATTTCAG
1760	1780	*	1800
GCAATCTGCA GACTTAGGAT	TGGATGGCGT	TCAGGAGCTT	GGATTGGTT TCTCACATCA
1820	1840		1860
TATTATTA ATAATTATT	AATTAAATT	TATGGACTTT	TGGACTGTCT GACTAATT

Figure 2C

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1880	1900	1920	
CAGAATTAA TTTTGGTTT GGGTTTTGTT GAATTTTTA GATAATTATT TTAAATTC	*		
1940	1960	1980	
TGCATTAATT TTCTGTTATT TGAAAAGGAT GTTCGAATT TTTTCAAAA TTGAAACGTT			
2000	2020	2040	
TAAGAATTAA TACTACTGCA AATTCAAAAT AAGTGAATT GTTTTTTAGA AAGATTAAT			
2060	2080	2100	*
AAGTTAGTAT TAGGATTAA AGTTTGATT GGTGGAAAGT AATGTATGTT TTTGAACATA			
2120	2140	2160	
ATTATTGAC ATAATTAAG TTTCAGG AATAAACCGGA AATATCTCT TCTTTTTGTT			
2180	2200	2220	
AAAATTACTA ATGCAAGAAC AAACAACGTT TTGGGGAGCA AATAATCTAG CTTTAAGTAG	*		
2240	2260	2280	
TCAGTGTAAAC TCTCAAAATC TGGTCATAAC TTCTAGGCTG AGTTTGCTGT GCTACAGTAG			
2300	2320	2340	
TAAGTCTATA GAAACTTACC TGACAAAACG ACATGACGTC AGGGTCGAAT CTACAACTTT	*		
2360	2380	2400	*
TCCTTTTCT TCAATTAAACA TATGGTTGAT TCAAGTCCG ATCTATAATA ATTATATTACG			
2420	2440	2460	
ATTATCAAT TTCAATTACC TTATATCATC CTATTATAAA TATAAGTCAG TTCAATTACAG			

Figure 2D

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2480	TTTCGAAAG TTCCCAAAA TTTCGAATT TATTAATT TTTCCTAAA ACCGAAATAG	2500	*	2520
2540	TTATATCTT CAAATTAG TTTCATTCTT CAATCCGATT TCAATTCTAT CCTTTTATAA	2560		2580
2600	*	2620		2640
2660	CTCTCTTAA TCTATAATTA CATAAATTTC AAATTAAATT TGAAATATT ACACTTAGT	2680	*	2700
2720	CCCTAAGTTC AAAACTATAA ATTTCACTT TAGAAATTAA TCATTCTCA CATCTAACCA	2740		2760
2780	TCAAATTAA CCAATGACA CAAAATTCTAT GATTAGTTAG ATCAAGCTTT TGACTCTTCA	2800	*	2820
2840	AAACATAAAA ATTACAAAAA AAAAACAAAC TTAAATTCTAT TTATCAATT TTAAACACAAA	2860		2880
2900	GCTTGGCGA ATGCTAAGAG CTAAAATG GCTTCCTTTTG TTTCCTTTTG TTGCAAACGG	2920		2940
2960	TGGAGAGAAG AGGGAAATGA AGATTCGACCA TATTCTTTA TTATGTTTA ACATATAATA	2980		3000
3020	TTAATATT ATTCAATT ATACTTTGGT GAATGTGACA GTGGGGAGAT ACGTAAAGTA	3040	*	3060
3080	TTTAACATT ATACCTTTG CAAGCAGTTG GCTGGTCTAC CCAAGAGTGA TCAAAGTTG	3100		3120

Figure 2E

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AGCTGCCTTC AATGAGCCAA	TTTTTGCCCC TAATGGATAA	AGGCAATTG TTTAGTTCAA	*
3140	3160	3180	
CTGCTCACAG AATAATGTTA	AAAATAATT AAAATAAGGT	GGCCTGGTCA CACACACAA	
3200	3220	3240	
*			
AAAAAACTAA TGTTGGTGG	TTGAATTATA TATTACGGAA	TGTAATAATT TATTTTAAAA	
3260	3280	3300	*
TAAAATTATG TTATTTAGAT	TCTTAATATT TTGGGAGCATT	CCATACTATA ATTTCGTAAC	
3320	3340	3360	
ATAAATTTAA AATATAGTAA	TATAAAGTGT AATTAACCTT	AAATTACAAG CATAATATTA	
3380	3400	3420	*
AATTTGAAAT CAATTAAATT	TTTATTCTAT TATTTTAATT	AAATTAGTCT ATTTTTTCAA	
3440	3460	3480	
AATAAAATT AAATCTAAAT	AAAATAATT TTTCCCTTAAT	GTTGAAACCAA CTCATGTTAT	
3500	3520	3540	
*			
ACTTCAAAAT TATAAAGTATT	ATATTACCT TGATGATTAA	TTTATTAGTA TATTAATTCT	
3560	3580	3600	*
GATTATAATT ATGGTGGGAT	ACAATCGCTT TCCACTAAAT	TTTTAACTA TGATTATAA	
3620	3640	3660	
ATTTTATTCA ACATCGTATA	TTTACTTATT AATAACATAAT	TTATCATAAT TTTATGGAAA	
3680	3700	3720	*

Figure 2F

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TTGAGACCAA GAAACATTAA GAGAACAAAT TCTATAACAA AGACAATTAA GAAAAAATG  
 3740 3760 3780  
 TACTTTAGG TAATTTAGG TACTCTTAAC CAAACACAAA ATTCAAAATC AAATGAACTA  
 3800 \* 3820 3840  
 AATAAGATAA TATAACATAC GGAACATCTT ACTTGAAATC TTACATTCCC ATAATTTTAT  
 3860 3880 3900 \*  
 TATGAAAAAT AATCTTATAT TACTCGAACT AAATGTGTC ACAAAATTAT ATCTAAATAA  
 3920 3940 3960  
 AGAAAAACAC TTAATTTTA TAACATTTT TCATATATT GAAAGATTAT ATTGTGATA  
 3980 4000 4020  
 \*  
 TTTACGTAAA AATATTGAC ATAGATTGAG CACCTTCTTA ACATAATCCC ACCATAAGTC  
 4040 4060 4080  
 AAGTATGTAG ATGAGAAATT GGTACAAACA ACGTGGGGCC AAATCCACC AAACCATCTC  
 4100 \* 4120  
 \*  
 TCATTCTCTC CTATAAAAGG CTTGCTACAC ATAGACAAACA ATCCACACA C AAA TAC  
 <Phe Val  
 4140 4160 4180  
 ACG TTG TTT TCT TTC TAT TTG ATT AAC CAT GGC TCA TAG CAT TCG TCA  
 <Arg Glu Lys Arg Glu Ile Gln Asn Val Met Ala \*\*\* Leu Met Arg \*\*\*  
 4200 \* 4220  
 CCC TTT CCT CCT TTT CCA ACT TTT ACT CAT AAG TGT CTC ACT AGT GAC  
 <GLY Lys Lys Arg Lys Trp Ser Lys Ser Met Leu Thr Glu Ser Thr Val

Figure 2G

4240	CGG TAG CCA CAC TGT TTC GGC AGC GGC TCG ACG TTT ATT CGA GAC ACA	4260
<Pro Leu Trp Val Thr Glu Ala Ala Ala Arg Arg Lys Asn Ser Val Cys		
4300		4320
	*	
4340	GAG AGT CTG AAT ACG AAA AGC CAG AAT ACA AAC AGC CAA AGT ATC ACG	4360
<Leu Thr Gln Ile Arg Phe Ala Leu Ile Cys Val Ala Glu Phe Tyr Ser Leu Val		
4380		4400
	*	
4440	AAG AGT ACT CAA AAC TTG AGA AGC CTG AAA TGC AAA AGG AGG AAA AAC	4460
<Leu Thr Ser Leu Val Gln Ser Ala Gln Phe Ala Phe Pro Pro Phe Val		
4480		4500
	*	
4540	AAA AAC CCT GCA AAC AGC ATG AAG AGT ACC ACG AGT CAC ACG AAT CAA	4560
<Phe Val Arg Cys Val Ala His Leu Thr Gly Arg Thr Val Arg Ile Leu		
4660		4680
	*	
4700	TGGTGATTGG GAAATGTGTG TGTGCAATTCC TCCATGCACT AATGGTGAAT CTCTTTGCAT	4720
	*	

Figure 2H

ACATAGAAAT	TCTAAATGTT	TATAGTTAT	GTTATAGTGT	ATGTTGTAGT	GAATTAAATT	4720	4740	4760
4780		4800	*			4820		
TTAAATGTTG	TATCTAATGTT	TAACATCACT	TGGCTTGATT	TATGTTATGT	TATGTTATT	4840	4860	4880
ACTTTAATGAA	TATTGCATGT	ATTGTTAATT	TAACATGCT	TGATCATTAT	ACTCTTCCTAC	4900	4920	4940
*								
TATTAATTAT	AAATGGCACT	GTTTGTGTTA	AACCTTTTAC	AAAGTTAAGGAC	ATGTATAAAT	4960	4980	5000
			*					
ATATGACAAT	ATAATTACAG	GTTTTAGTTC	AATGTTAGCT	ATCTTAGTAT	GTTTATTGATG	5020	5040	5060
ATCTTAATTA	CATTAAACA	AATTCCACTT	AAAATTAA	TAATAATAAA	CAAATAATTA	5080	5100	5120
			*					
TTGTAATATA	ATACATTAA	TGCAACAAAA	AATGAAATAA	ATAAAATAAA	ATAGCAAATA	5140	5160	5180
ATGTTATAA	TATTGTATAA	TAATATGTCAC	CATATTCTTA	ACTGAAATAG	GGTCTAACCT	5200	5220	5240
*								
ATAATCCCTA	AAATTTCAGT	TTAAATATTT	TTATACCTAC	CATATTATA	GAACTCTTT	5260	5280	5300
					*			
TAAATATATT	AAAATTAA	TTATACCAAT	TTAATTAAC	TATTAATTAT	CTTAACTAAA			

Figure 2I

ATCTAAATT TTATTTAAC	5320	TATTAATAAA	TTCCTTAATTAA	TCTTATCTAA	TTTAAA	ACTC	5340
	5380	*	5400		5400		5420
TAATTATCCT AATTTAATT	5440	AAATTCTTAA	TTATCTTAAT	TGTAACCTC	CTCCACCCAG		
CTAGATGCTG GACCCGAATC	5440	CGGGAGATTAA	CATCGGCCAT	TGAGATGGCG	TGATCAGGGT		5460
	5500	*	5520		5540		5480
TTGGCGGCC GGTACCCAAT		TCGCCCTATA	G TGAGTCGT	ATTACGGCG	CTCACTGCGT		
			CCGGTTT				

Figure 2J

ACTAAAGGGA ACAAAAGCTG 20 GAGCTCCACC 40 GCGGTGGGG CCGCTCTAGG ATCCCCCGTG  
 GACTAAACAA AACATGGGA GATTGCTGT 80 100 \* 120  
 CTTTGTGAAT TGTATAACAA 140 AGACTCAATG 160 AAAAACATA ACTCAATACA 180  
 200 220 240  
 \*  
 CTGATTTACA TCCTTTATAT AGGCTGAAC TACAAACACT TTAGCTAAAAA AAATAGGATA  
 260 280 300 \*  
 ACCTAAATAGC AAAATCACAA TCAGATATTA AACCATGATT TTAGCTAAC 300  
 TTTATTGAAA CTAATTGAA TATTTCATCT GCTGATATGC 320 340 360  
 CCAAGATTIT AGGCCACTAA  
 380 400 \* 420  
 CCGATTGGT GGTGAACTT AACATGTCAT GCATTGTAA CTGTTGAAA CAAGTTTTT  
 GCATTATTTT ACTATATGAA CTGTTGATT AGGTGAGTT ACACACTGAG CTTGTAAGCT  
 440 460 480  
 CACTCAAATT TTCTAAATT CTAAGGTGAT CAGCAAACCTT AGGACCGGGC GGCCTACGAG  
 500 520 540  
 AGCTCGGATT GATTTCCTAG TTAATAATAAAGACGATTAA TGTTTTAACTATTATGGA  
 560 580 600 \*  
 \*

Figure 3A

CTTTTGGAC TATGTAAC TG	620	TTTGGACTT TATTGTT	640	TTTTTGC TTGGA	660
	680	*	700		720
TTAGTAAATT ATTATTTTA AACTGCCAAA TTATGTT		TTACAAACTA AGTCACAGTT			
TTCAAATT CATAACTTAG AATTTCAGC	740	TGCAAAATAA AGTAATCATT TAAGTGT	760		780
	800	*	820		840
TTCTGTAAATA AAATAAAATAA ATAATTAA CGAGTATT CCTAAAAATT GGAAATTGAT					
	860		880		900
TTACCAAAAT TAGTATGTC AAACACATGT TTATGTTA CAGGGCGATA TCGTCTAGGC		*			
AAATAACATC TAGGGGGGT TTGGAGTGT ACAGGGGAG TGGGCTCATT TTGAGTAAGT	920	940	960		
	980	*	1000		1020
ATAGTTAGGG CCGAGTTTA GATTGCATAT TCAAGGTCAA AGATTTTGTA AACTTCGATG					
AATGATATGAT ATGATTGTCC GATTAACGAA ATATGTTTT TCTTTGTG TGTGTTTAT	1040	1060	1080		
	1100	*	1120		1140
CTCGTGTGAT AAGTATATAG TATGTTTAT TCCAATTCTT ATGGCATGTG ACATGTGGC					
TATTCATATT AAATTGATT GTTATTATG AAATCTGATG CATCTGTTCT ACMAAGCATG	1160		1180		*
	1220		1240		1260

Figure 3B

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GAATCTCATG	CCTACTGGCT	TCTGTTAAAG	ATACGATTGC	AAGTTAACCA	TGCTTACTAT
1280		1300		1320	
TTTGATTGG	TCCTTGCATG	CTATGTCACA	TTACATGGGG	TTGGGATGAT	ATGGTAAGGA
1340		1360		1380	
GCAAGTTTG	ACAGTTAAT	GATTGCACT	ATCTGGTGGT	TTAACACAT	ATTGGTTATG
1400	*	1420		1440	
GCATCTTGAC	TGGCGTTATG	GTGGCTCGAC	CGCCCATATC	TGTTCTGGAA	ATTATCTGT
1460		1480		1500	*
GAECTCTGGTG	GCATTGCTA	CAATTATTG	TTGGTGTGTT	TTGGATGGAC	GAGTCGTGGG
1520		1540		1560	
GAACCTATT	TGGTGTGTTG	CGGAGTTGGG	TAGGAAATT	TCGAAAAAAA	TTTGCATTGT
1580		1600	*	1620	
GTTRTTCTGA	AAAATATTGC	ATTAACATAA	TCATGCATT	TCAATTGTTGG	TCAATTGAAAC
1640		1660		1680	
GTATAAAAT	TCTCTATGAT	ATCCCTGATCT	GTGTTAACCA	TTATATGTT	TTATGCTTGA
1700	*	1720		1740	
GTAAAGTCAA	ACATTGAGAT	TCATAGCTCA	CCCAATTATT	TAATCATTTC	AGGCAAATCTG
1760		1780		1800	*
CAGACTTAGG	ATTGGATGGC	GTTCAGGAGC	TTGGATTGGT	TTTCTCACAT	CATATTAT
1820		1840		1860	
TAATAATTAA	TTAATTAAA	TTTATGGACT	TTGGACTGT	CTGACTAATT	TTCCAGAATT

**Figure 3C**

1880	1900	*	1920
TATTTGGTT TTGGGTTTG TTGAAATT TTGATAAATTAA TTTTAATAT TCTGCCATAAT			
1940	1960	*	1980
TTTTCGTAA TTGAAAAGG ATGTTCGAAT TTTTTTCAA ATTGAAACG TTAAAGAATT			
2000	2020		2040
*			
TTTACTACTG CAAATTGAGA ATAAGTGAAT TTGTTTTTA GAAGATTAA ATAAGTTAGT			
2060	2080		2100
ATTACGATT TTAGTTGAT TTGGTGGAAA GTAATGTATG TTTTGAAACA TAATTATTG		*	
2120	2140		2160
ACAATAATTAA AGTTTTCTAG GGAAATAAACG GAAATATCTT CTTCCTTTTT GTAAAAATTAC			
2180	2200		2220
TAATGCAAGA ACAACAAACG TTTCGGGAG CAAATAATCT AGCTTTAAGT AGTCAGTGTAA	*		
2240	2260		2280
ACTCTCAAAA TCTGGTCATA ACTTCTAGGC TGAGTTTGCT GTGCTACAGT AGTAAGTCTA			
2300	2320		2340
*			
TAGAAACTTA CCTGACAAAA CGACATGACG TCAGGGTCGA ATCTACAAC TTTCTTTT			
2360	2380		2400
CTTCATTAA CATATGGTG ATTCAAGTTC CGATCTATAA TAATTATA CGATTATCA	*		
2420	2440		2460
ATTTCATTAA CCTTATATCA TCCTATTATA AATATAAGTC AGTTCAATTG AGTTTCGAA			

Figure 3D

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2480	2500	2520
AGTTCCAAA AATTGGAT TTTATAAT TTATTCCCTA AAACCGAAAT AGTTATATCT	*	
2540	2560	2580
TTCAAAATTAA AGTTTCATT TTCAATCCGA TTTCAAATTTC ATCCTTTAT AACTCTCTAT		
2600	2620	2640
TATCTATAAT TACATAAATT TCAAATTAAAT TTTGAAATAT TTACACTTA GTCCCTAAGT		
2660	2680	2700
TCAAAACTAT AAATTTCAC TTTAGAAATT AATCATTTT CACATCTAAG CATCAAATT	*	
2720	2740	2760
AACCAAATGA CACAAATTTC ATGATTAGTT AGATCAAGCT TTTGAGTCTT CAAACATAA		
2780	2800	2820
AAATTACAAA AAAAAACAA ACTTAAAAATC ATTATCAAT TTGAACAAACA AAGCTGGCC	*	
2840	2860	2880
GAATGCTAAG AGCTTAAAAA TGGCTTCCTT TGTTTCTTT TGTTGCAAAC GGTGGAGAGA		
2900	2920	2940
AGAGGGAAAT GAAGATGGAC CATATTTT TATTATGTT TAACATATAA TATTAATAAT		
2960	2980	3000
TTAATCATAA TTATACCTTG GTGAATGTGA CAGTGGGAG ATACGTAAG TATTAAACA	*	
3020	3040	3060
TTTATACCTT TGCAAGCAGT TGGCTGGCT ACCCAAAGAGT GATCAAAGTT TGAGCTGCCT		
3080	3100	3120

Figure 3E

TCAATGAGCC	AATTTTGGCC	CATAATGGAT	AAAGGCAATT	TGTTTAGTTC	AACTGCTCAC	*
3140		3160		3180		
AGAATAATGT	AAAATGAAA	TTAAAATAAG	GTGGCCTGGT	CACACACACA	AAAAAAACT	
3200	*	3220		3240		
AATGTTGGTT	GCTTGAATT	TATATTACGG	AATGTAATAT	TATATTCAA	AATAAAATTA	
3260		3280		3300		*
TGTTATTAG	ATTCTTAATA	TTTGGAGCA	TTCCCATACTA	TAATTTCGTA	ACATAATATT	
3320		3340		3360		*
AAAATATAGT	AATATAAAGT	GTAATTAACT	TTAAATTACA	AGCATAATAT	TAAATTGTA	
3380	*	3400	*	3420		
ATCAATTAAAT	TTTTTATTCT	ATTATTCAA	TTAATTAGT	CTATTTC	AAAATAAAAT	
3440		3460		3480		*
TTAAATCTAA	ATAAAAATAAA	TTTTTCCTTA	ATGTTGAAAC	AACTCATGTT	ATACTTCAAA	
3500	*	3520		3540		
ATTATAAGTA	TTATATTAC	CTTGATGATT	TATTTATTAG	TATATTAAATT	CTGATTATAA	
3560		3580		3600		*
TTATGGGG	ATACAATCGC	TTTCCACTAA	ATATTTAAC	TATGATTAT	AAATTATTT	
3620		3640		3660		*
CAACATCGTA	TATTTACTTA	TTAATACATA	ATTTTATCATA	ATTTTATGGA	AATTGAGACC	
3680		3700	*	3720		

Figure 3F

AAGAAACATT AAGAGAACAA ATTCTATAAC AAAGACAAATT TAGAAAAAA TGTACTTTA  
 3740 3760 3780  
 GGTAAATTTA AGTACTCTTA ACCAACACAA AAAATTCAA TCAAATGAAAC TAAATAAGAT  
 3800 \* 3820 3840  
 AATATAACAT ACGGAACATC TTACTTGAA TCTTACATT CCATAATT ATTATGAAA  
 3860 3880 3900 \*  
 ATAATCTTAT ATTACTCGAA CTAATGTTG TCACAAATT TAATCTAAAT AAAGAAAAAC  
 3920 3940 3960  
 ACTTAATTTT TATAACATT TTTCATATAT TTGAAAGATT ATATTTGTA TATTACGTA  
 3980 4000 \* 4020  
 AAAATATTG ACATAGATG AGCACCTTCT TAACATAATC CCACCATAG TCAAGTATGT  
 4040 4060 4080  
 AGATGAGAAA TTGGTACAAA CAACGTGGGG CCAAATCCC CCAAACCATC TCTCATTCTC  
 4100 \* 4120  
 TCCTATAAAA GGCTTGCTAC ACATAGACAA CAATCCACAC A CA AAT ACA CGT TCT  
 < Ile Cys Thr Arg  
 4140 4160 4180  
 TTT CTG TCT ATT TGA TTA ACC ATG G CTCATAGCAT TCGTCACCCCT TTCTTCCTTT  
 < Lys Arg Asn Ser \*\*\* Gly His  
 4200 \* 4220 4240  
 TCCAACCTTT ACTCATTAAGT GTCCTCACTAG TGACCGGTAG CCACACTGTT TCGGCAGCGG  
 4260 4280 4300

Figure 3G

CTCGACGTTT ATT CGAGACA CAAGCAACCT CATCAGGCT CCCACAATTG GCTTCAAAAT  
 4320 4340 4360  
 ACGAAAGCA CGAAGAGTCT GAATACGAA AGCCAGAATA CAAACAGCCA AAGTATCACG  
 4380 4400 4420  
 \*  
 AAGAGTACTC AAAACTTGAG AAGCCTGAA TGCAAAGGA GGAAAAACAA AAACCCCTGCA  
 4440 4460 4480  
 AACAGCATGA AGAGTACAC GAGTCACACG AATCAAAGGA GCAAAAAGAG TACGAGAAAG  
 4500 4520 4540  
 \*  
 AAAATCTCGA CGGGCCCCGAA GATCTTCGCT AGCCGTGAC GCCCGGGGA ATTCTGCGAG  
 4560 4580 4600  
 \*  
 CCTTGATCA TATGACGGCTG GTGCATGTGC CATCATCATG CAGTAATTTC ATGGTATATC  
 4620 4640 4660  
 GTAATATAA GTTAATAAAA AAGATGGTGA TTGGGAAATG TGTGTGTGCA TTCCCTCCATG  
 4680 4700 4720  
 \*  
 CACTAATGGT GAATCTCTTT GCATACATAG AAATTCTAAA TGTTTATAGT TTATGTTATA  
 4740 4760 4780  
 GTGTATGTTG TAGTGAAGKT AATTTTAAAT GTTGTATCTA ATGTTAACAT CACTGGCTT  
 4800 4820 4840  
 \*  
 GATTATGTT ATGTTATGTA TTTTACTTTA ATGATATTGC ATGTTATGTT AATTAAACAT  
 4860 4880 4900  
 \*

Figure 3H

TGCTTGATCA TTATACTCTT CTACTATTAA TTATAATGG CACTGTTTG TTAAACTTT  
 4920 4940 4960  
 TTACAAAGTTA AGACATGTAT AAATATATGA CAATATAATT ACAAGTTTA GTTCAATGTT  
 4980 5000 \* 5020  
 AGCTATCTTA GTATGTTATT GATGATCTTA ATTACATTTA ACAAAATTCC ACTTAAATT  
 5040 5060 5080  
 TTAATAATA ATAACAATA ATTATGTAA TATAATACAT TAAATGCCAAC AAAAATGAA  
 5100 5120 5140  
 \* ATAAATAAA TAAAATGCCA AATAATTGTT ATAATATTGT AATAATAATTAT GTACCCATT  
 5160 5180 5200 \*  
 CTTAACTGAA ATAGGGTCTA ACCTATAATC CCTAAAATT CAGTTAAAT ATTTCATAC  
 5220 5240 5260  
 CTGCCCATATT ATTAGAACCTC TTTTTAAATA TATTAATAATT TTAATTATAAC CAATTAAATT  
 5280 5300 \* 5320  
 TAAACTATTAA ATTATCTTAA CTAAAAATCTA AAATTATTATT TAACCTATT ATTAAATTCC  
 5340 5360 5380  
 TAATTATCTT ATCTAATTAA AACACTCTAAT TATCCTTAATT TGATTAAAT TCTTGATTAT  
 5400 5420 5440  
 \* CTTAAATTGTT AACCTCCCTCC ACCCAGCTAG ATGCTGGACCC CGAATCCGGG AGATTACATC  
 5460 5480 5500 \*  
 GGCATTGAGA TGGCCTAGTA GTGATCAGGG TTTCTAGAG GTACCCAATT CGCCCTATAG

Figure 3I

TGAGTCGT

Figure 3J

AAAAACA ATG AGC ACT GCA AGA TTT ATC AAG TGT GTC ACG GTC GGT GAT	50
Met Ser Thr Ala Arg Phe Ile Lys Cys Val Thr Val Gly Asp	
1	
GGA GCT GTG GGG AAA ACT TGT ATG CTC ATT TCA TAT ACC AGC AAT ACT	98
Gly Ala Val Gly Lys Thr Cys Met Leu Ile Ser Tyr Thr Ser Asn Thr	
15	20
TTC CCA ACG GAT TAT GTT CCA ACA GTA TTT GAT AAC TTT AGT GCC AAT	146
Phe Pro Thr Asp Tyr Val Pro Thr Val Phe Asp Asn Phe Ser Ala Asn	
35	40
GTG GTG GAT GGC AGC ACA GTG AAC CTT GGC CTA TGG GAC ACT GCC	194
Val Val Val Asp Gly Ser Thr Val Asn Leu Gly Leu Trp Asp Thr Ala	
50	55
GGG CAA GAA GAT TAT AAT AGG CTA AGG CCA CTG AGT TAT AGA GGA GCT	242
Gly Gln Glu Asp Tyr Asn Arg Leu Arg Pro Leu Ser Tyr Arg Gly Ala	
65	70
GAT GTG TTT TTG TTG GCC TTT TCT CTT ATA AGC AAG GCC AGT TAT GAA	290
Asp Val Phe Leu Leu Ala Phe Ser Leu Ile Ser Lys Ala Ser Tyr Glu	
80	85
AAC ATC TAC AAA AAG TGG ATC CCA GAG CTA AGA CAT TAT GCT CAT AAT	338
Asn Ile Tyr Lys Lys Trp Ile Pro Glu Leu Arg His Tyr Ala His Asn	
95	100
GTA CCA GTT GTG CRT GTT GGA ACC AAA CTA GAT TTG CGA GAT GAC AAG	386
Val Pro Val Val Gly Thr Lys Leu Asp Leu Arg Asp Asp Lys	
115	120
CAG TTC CTC ATT GAT CAC CCT GGA GCA ACA CCA ATA TCA ACA TCT CAG	434
Gln Phe Leu Ile Asp His Pro Gly Ala Thr Pro Ile Ser Thr Ser Gln	
130	135
GGA GAA GAA CTA AAG AAG ATG ATA GGA GCA GTT ACT TAT ATA GAA TGC	482

FIGURE 4A

Gly	Glu	Leu	Lys	Lys	Met	Ile	Gly	Ala	Val	Thr	Tyr	Ile	Glu	Cys		
145																
AGC	TCC	AAA	ACC	CAA	CAG	AAT	GTG	AAG	GCT	GTT	TTC	GAT	GCT	GCA	ATA	530
Ser	Ser	Lys	Thr	Gln	Gln	Asn	Val	Lys	Ala	Val	Phe	Asp	Ala	Ala	Ile	
160																
AAA	GTA	GCT	TTG	AGG	CCA	AAA	CCA	AAG	AGA	AAG	CCT	TGC	AAA	AGG	578	
Lys	Val	Ala	Leu	Arg	Pro	Pro	Pro	Lys	Pro	Lys	Arg	Lys	Pro	Cys	Lys	Arg
175																
AGA	ACA	TGT	GCT	TTC	CTT	TGA	ATATTGG	ATC	ATTATA	CAGTC	AAAAA					626
Arg	Thr	Cys	Ala	Phe	Leu											
180																
CAGTTAACAA	AAGCTGTGCA	AGATAAACAC	TGAATCTGCT	ATAGTTGTT	TTTGGTTTAC											686
ATATGTTCCA	CGTGAAACTA	TGAAGCATCT	CTAAGAAAC	CCAAACTATC	ATATCAACCC											
ATCGATCAAT	GAATCGATT	CAATTTCGGC	AGTATAAGTT	CCTTCTTAATC	CTTTCCTTFT											
ACTTCATT	ATAACGAATT	CTATGGATAA	TGTTCCCTAC	AAACATGTCA	TTACAATGT											
TAATTATAAA	TTCCATTCTT	CTATTTACT	AAAAAA	AAAAA	AAAAA											
															910	

FIGURE 4B

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20                    TTGGATGAGA ACCAATTTT AATAGTAAAN CCTAACCAAT TTTAATAAT AAAGCTGACT  
 5                    80                    100                    \*  
 CCTAGTACAA GAGCTTTAT TCATTCTCT ATTTCCTTT CCTCTAGGCT TGGCAATCGA  
 10                    140                    160                    180  
 GAATTTCCT GTGTTACAT ATAATAAATA CATCGTAA ATAATTAA TTCAAATTGA  
 20                    200                    220                    240  
 AGTCCTAACC ATCTTTAATA TTTCCTAGATG TAATTTAAAT GAAAGATAAA TACATATTCT  
 15                    260                    280                    300                    \*  
 TGGACATGTA TTTTCATCTT AATGTTTGTG GCTTGTGTA TAGGTGTATT GATGTACGAT  
 20                    320                    340                    360  
 GTCTTTAAA TCACATATCA CATTTTGAGT TTGTATGATG ATAAGTCGAC ATANNGAMA  
 25                    380                    400                    420                    \*  
 TATGGTGTGA TCTTCACCTT TGAACTTGA TAAGTCACCA AACTTAACCA AAGTTTGATT  
 30                    440                    460                    480  
 GTGTACATAT ATATATATAT CTTCAAAATT TATAATAAAA ATGTGTATA AATAATTAC  
 35                    500                    520                    540                    \*  
 AGTTATATTA TTTTTTATC TCTAATTATA TTGTGCCA AATTTTAGT TGATATTAA  
 35                    560                    580                    600                    \*  
 ACATAAAAAA AATTGTACAC ATTACAAGC CCATATACAA ATAATTATAT AAATATTCA

FIGURE 5/A

5

TAAAAAT ATTAAAT	620	AGGATAATAA	TATAACTATT	640	TAGAATTAT	TCTACTTAA	660
	680	*		700			720
GATAACATAG GTAAATGTA	TAATTAATAA	GGTTAGTTA	TGTAAAGAT	GAGTATATAT			
	740	CCATTTTAT	TAACTCTTG	GTTCAGT	TCCAAAAGA	780	
	800	*		820			840
AAATGGAGG GAAATTGAG	AGTAAGTTCA	TGTTTATATT	ATACATAATG	AAGTGTATGT			
	860	880		900	*		
TTTCTCTTT TTAATATT	TATCAAAT	ATTAAATAA	AATAATTAG	GATTGAATGA			
	920	TTACTAATAG	TCATATGCA	TTCAGCGCA	TCTACTTAA	960	
AAAATATAAT GAAAGTCGTT	TTACTAATAG	TCATATGCA	TTCAGCGCA	TCTACTTAA			
	980	*	1000	*		1020	
TAATAGATAA ATTAATTGTG	GTACATTAGA	TCAAAGAAC	AACTAGATT	TGTCCCATTC			
	1040	TITACATTAA	AATAAGGTAC	ATGTTACATG	CCACGTATAA	1080	
	1100	*	1120			1140	
CTATCTGGTT ATTCTATCAA	TCACCGCTAAT	TTTAACAGT	AGAAATGAAT	GTAATTTTTA			
	1160		1180			1200	*
AATAGAAAGG GTCAAATTGT	TATTGATCT	AACACGTAGG	GATTAATTAA	CTTAACTTCC			
	1220		1240			1260	

FIGURE 5/B

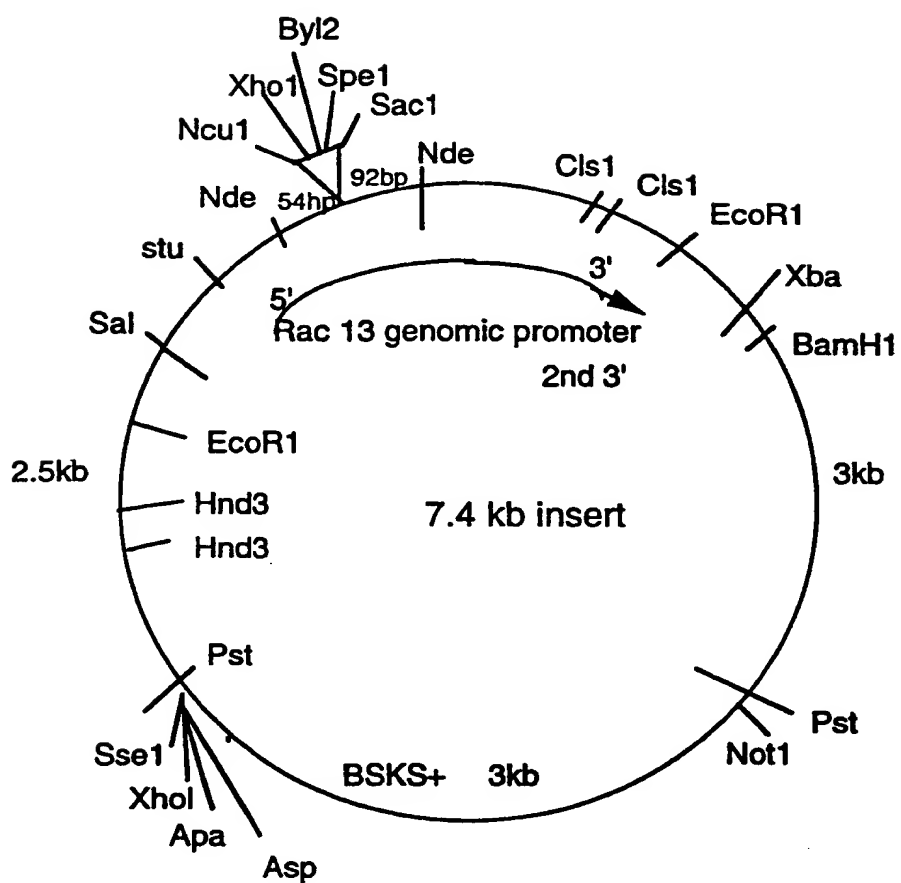
TAAAGAAATA AGTAAAATATT AATTGAATC TTAATACAAA AACTTCATG ATACTTTAT  
 1280 1300 \* 1320  
 5 CATACTTAC TTATAATTAA ATTATGTGAG AGTAACAAAR TTAAAAACA TAGAAACACC  
 AAAAGTTAGT TATGGTGTGA CTCATATAACA CAGTTAAAT TTGAATAAT TTTTTCTTC  
 1340 1360 1380  
 10 1400 1420 1440  
 GTCAATTAAATT CCATCATGGG TTTTTTTT TCTAGTTAAG CCATAATTAT CAAATAATC  
 1460 1480 1500 \*  
 15 ATCAATTAAATC CTATCAAATAC CCCGCCCTGC CTCCCTCCCT CAATACTTAA ACCAACTAA  
 CACCCAGCAC CAAACGCAC TTAATAGCCA CCTATTCTA GCATGTCCCT TGCACTTAA  
 1520 1540 1560 \*  
 20 1580 1600 \* 1620  
 GAAAAGTAAA GCTAACCTGCA AATCATTCCA TATCGAGGCC TCAACAGATA AAGTTGGTGTG  
 1640 1660 1680  
 25 ATGGGTTTGC ACCAAGTTGT TAAAACCCGG CCCTCAACTT CCCTTTCTT TTCATCCTCC  
 1700 1720 1740  
 30 CCACTCCACA CCCTCCAATT TTCTTCATAT GGTTCTTAA TAAGTTCTT ATAATCACAG  
 1760 1780 1800 \*  
 35 AATCAAGATA AGTCCTCAGC AAACAAAAAA CCATGGCTCT CGAGCAAGAT CTGGACTAGT  
 1820 1840 1860  
 CAGAGGCTCTG AATATTGGAT CATTATTACA GTCAAAACAA GTAAACAAAA GCTGTTGCAG

5		1880	1900	1920
	ATAAACACTG AATCTGCTAT AGTTTGTCTT TGTTTACAT ATGTTCCACG TGAACATATG	*		
	1940	1960	1980	
	AAGCATCTCT AAGAAACCC AAACTATCAT ATCAACCCAT CGATCAATGA ATCGATTCTA			
10		2000	2020	2040
	ATTTTCGGCAG TATAAGTTCC TTTTAATCCT TTCTTTTAC TTCATTAT AACGAATTCT	*		
	2060	2080	2100	
	ATGGATAATG TICCCCTACAA ACATGTCATT ACAATGGTTA ATTATAAATT CCATTCTCT	*		
15		2120	2140	2160
	ATTTTACTAA GATATTAGTA ACTTCAAACT GCTGATTCTT ACTAATTAT TATTATAAA			
	2180	2200	2220	
	TGTTAGAAT GATTATTCTT CAATAATTAA ACAACAAAT TTAATATTAT TATTATTATT	*		
20		2240	2260	2280
	ATTTCTCAAT TTTTATTAAA CAAAAACATA AATTITGAC AAATTAAAT AAATGAATTA			
	2300	2320	2340	
	ATTTCTCAAT TTTTCGTGCA ACTATTACAA AAATCCCTCA TAGTCCTAAT CTTAATTGTA	*		
25		2360	2380	2400
	TGCAGAGGTG ATAATAATCT TAATTGATG CAGAGGTAAT AATGGGCCGG GTTGAGGCTG	*		
	2420	2440	2460	
	GACTTAAGCA TGATATTGAC GTACTTATAA TTTTCACAA TTCAACCCAG CTCGAAATAT			
30				
35				

FIGURE 5/D

	2480	2500	2520
		*	
5	2540	2560	2580
	TAATTAAAAA AATTATATAC ATTATTATT ATTATTAAAT TATTATTAT ATTTTTATT		
	2600	2620	2640
10	TATTGAAAAT TTTTATATAG TCATCTAAC ATTATGTTAA TGTGTTATATT AGAGTAGTAT		
	2660	2680	2700
		*	
15	TATATATTATT TAGTATAGGT TTATTTTGTT AATAAACTTA AAAATGGGTCA TTGTTGGGCTA		
	2720	2740	2760
	GACTTGGACC TAAATGCTC AAACCTCAAAC TAAATTCAA TTTAAACAG GCTTAATATT		
	2780	2800	2820
20	TTTATTACA CTGTTTCAA TTTTTCGGGT GAAATATCTT CGAGTCTAGA TTAATAAACAC		
	2840	2860	2880
	CACAGGTCTA ATTGTGATGCT CAATGAAAAT GAAATCATAT TGAGCTTAAT TAATATTCCA		
25	2900	2920	2940
		*	
	TTCCTCTTG CTGAAAGGAC CAAGCAATTG GAGTTACATT AAGGTTAAAG AGTATGGGAT		
30	2960	2980	3000
		*	
	CCGCCAAACC TGCCCCAATG TCTCTTCAAC CATCCAAAAA CTGAGTCAG TATCACATAC		
35	3020	3040	
	ATGTACCGNT ATTATTATT ATTATTCAAT TGGCATTTT TCTTIG		

FIGURE 5/E

**FIGURE 6**

GGGCATTCCA CACGACCAG TGTCCCTAT TTCCAGGCAT TTGAGACT CACCTAACT	60
TCTAGAGTTG TTTCAAATTA GCCCCTATT GTTCTTAAT CATTTAGGA TCTTGTAAC	120
TCGTATTAG GACTAAATGT GTAATTATA CTTTAATTAT GATGATTAA TGATTGATT	180
TNGTAGTAAT GCCCGTGAC CTAATCCGTT AGCGAAGAGG GTTAGGGCT TAGGGGTTT	240
TTTATTATT TTAGATATT GTATAACTCT TGTCTTATT TTAAATTGTT TACTATTCA	300
AAGGCATTG TTCTAGTGT TATTTCGAGT AGGTTTATG GGTGAACAAC CCTTGACCGC	360
CAAATCAATC ACAAGAGTTT AACATTTAT TTATTGAA ATGTATTAA AATCGTTAAT	420
CTATATATC GCCCCATTAT TGGGATTAAA TATTCAACAG GTTGTAGACC GTCATGAGAC	480
AGATTAGTTT TATCTTACTG ATGGTCACAT CACAATAGTA ATTCAACTTA ATACGAGAGG	540
AACCATTGAT TCACGCAATT GGTCATCGCA CTTAGTTGAA AAGCTAGGGG TGCGAAGCTA	600
CCGTACGGCTG GATTATGATT GAACACCTCT AAGTCAGAAT CGAAATTAGA AACAATGCCAC	660
GTGTCCGGTTG CCTGATTGCC AACCCCAATA ACACGTGTTG TAGTTTAAC CATGTTTATG	720
AAAGATAAGG TTTTTTTT TATAAGCAAG CAACTATAGG GTTGTACTTC CGTGGCAAA	780
TTTTTGGTT ACCTATTG GGAGGGGGAA TTATGATTCA AGTGAAGAA AGTTGGCACA	840
CACACAATCA GTACATCTGT TTTGACAGAG ACACAGCCTA AAAACAGCAG CAAACAGCC	900
TAAGGAAATC ACCAAAAAAC AACAAACAAA AGTACAGAGG AAAACAAAAG AATCCCTGTT	960
ACCAACCAAGC TGAAAAAAAG AAAATAAAAC TCAACTTTG GCAATAAAA CCCTCTTACCC	1,020
CTCAACCCCT AACACGGAA CAATCAGCAA TACTCCAGC AACCATTTC CTTACAAGTT	1,080

FIGURE 7A

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TGT TTTTCTT GTGATTAAAT CAT ATG GCT AGC TCC ATG TCC CTT AAG CTT GCA 1133  
 Met Ala Ser Ser Met Ser Leu Lys Leu Ala >

TGT CTG CTA GTG TTG TGC ATG GTC GTC GGT GCA CCC CTC GCT CAA GGG 1181  
 Cys Leu Val Leu Cys Met Val Val Gly Ala Pro Leu Ala Gln Gly >

GAC GTA ACC CGT GCT GAT GGC GTA GTC ACC CTT CCA CGC TGC CTT CCT 1229  
 Asp Val Thr Arg Ala Asp Gly Val Val Thr Leu Pro Arg Cys Leu Pro >

TTA TTG ATA GGG AAT GGT AAT GGT GCT GAT GCT GAT GTT GAT GCC CCA 1277  
 Leu Ile Gly Asn Gly Asn Gly Ala Asp Ala Asp Val Ala Asp Ala Pro >

GCT TGC TGC GAC ATC GTC AGG GGT CTC TTG AGC TCG CTG CTC TGT GGT 1325  
 Ala Cys Cys Asp Ile Val Arg Gly Leu Ser Ser Leu Leu Cys Gly >

GTT TAGAACCG ATCTAGCTTG AAATCGGGTT CGGATAACGGG TGGAGTTCA 1380  
 Gly Val >

AATTGGTGTG TTATGGAATC CCAAACTTAAT CGTGTAGG GGTGGGATCC AATTGGTGTGA 1440

TACATTACAG AGCATGGTTG TGGATGTTT TCTCATATGT TTGATTGAC TTGCTTGATA 1500

CATTGGATGA TTGATAAGG TGACCGGTTT ACCTGGGTAT CCAACCATCA TCCGATTACT 1560

TTTAATAAT TATTGGTTTC TTCTTTATGT TGTCTGTCTT TTGTTCTT GATCTATAAC 1620

ATTATATTG CCCAAATTTC CGCATTTTCC ATATGTAGCT TATATATGTA TATATATATT 1680

CAATAAAGTA TATGATTAA GCAGATGATT TGTGTATATA TTTAAATCAA ATCAAACATT 1740

ATGATCATT CACTAGCGTC TTAATCTTGA AAAATTCAATC AACGGTTATC CTTTGAGCCA 1800

TATATAAAA AAATTGCCAA CCCTATGCTT TTACACCTAA TTCAAGGGAT AACATAAGTC 1860

GATTAACG A 1871

FIGURE 7B

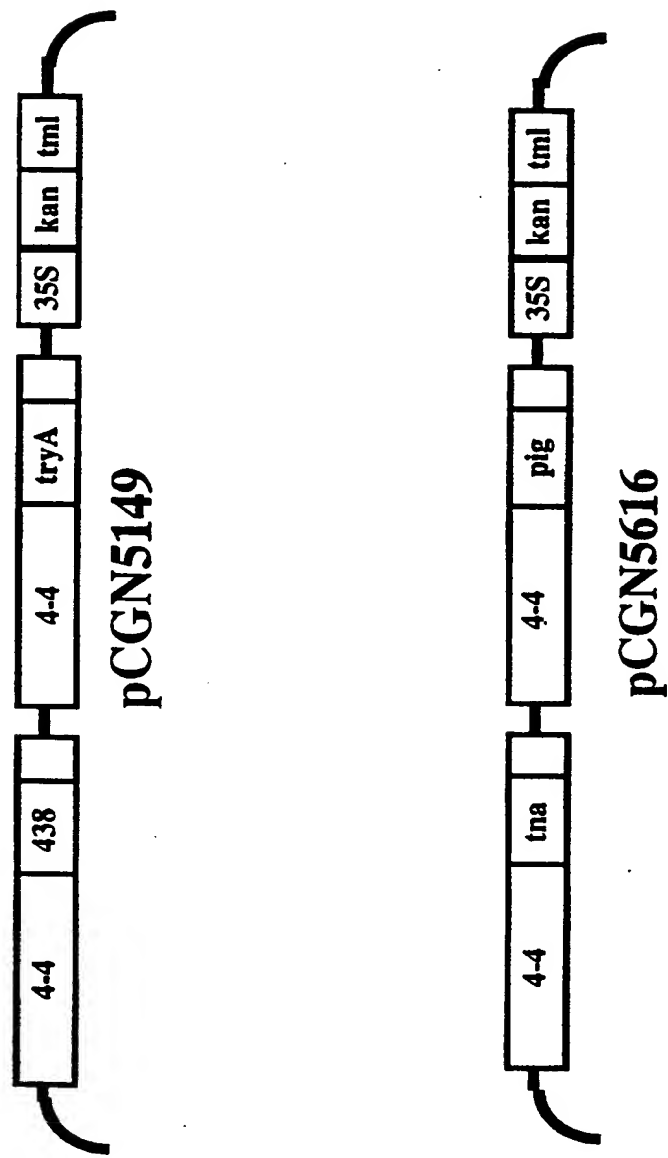


FIGURE 8

Coker 130	Y <sub>xy</sub> , Y	Y <sub>xy</sub> , x	Y <sub>xy</sub> , Y	Lab, L	Lab, a	Lab, b	LCh, L	LCh, C	LCh, h
1	80.35	.3206	0.3266	91.84	0.16	5.51	91.84	5.51	88.4
2	77.62	.3232	0.3282	90.6	0.68	6.45	90.6	6.48	84.2
3	80.98	.3197	0.3257	92.12	0.13	5.04	92.12	5.04	88.6
4	80.16	.3200	0.3255	91.75	0.35	5.00	91.75	5.01	86.1
5	77.03	.3220	0.3271	90.33	0.61	5.84	90.33	5.87	84.1
6	73.67	.3258	0.3293	88.76	1.35	7.14	88.76	7.26	79.4
7	82.43	.3178	0.3237	92.76	0.15	4.05	92.76	4.05	87.9
8	82.21	.3196	0.3255	92.66	0.19	4.99	92.66	4.99	87.9
9	81.19	.3194	0.3241	92.21	0.77	4.42	92.21	4.48	80.2
10	76.11	.3243	0.3229	89.9	0.74	6.89	89.9	6.92	84
11	82.28	.3178	0.3236	92.69	0.19	4.00	92.69	4.00	87.3
TOTAL	874.03	3.5302	3.5883	1005.62	5.30	59.33	1005.62	59.61	938.10
MEAN	79.46	.3209	.3262	91.42	0.48	5.39	91.42	5.42	85.28
S.D.	2.91	.0026	.0020	1.33	0.38	1.08	1.33	1.11	3.22
RANGE	82.43-73.67	3858.-3178	0.3293-.3236	92.76-88.76	1.35-13	7.14-4.00	92.76-88.76	7.26-4.00	88.6-79.4
AVER DEV.	2.44	.0021	.0017	1.11	0.31	0.88	1.11	0.90	2.64
<hr/>									
Coker 130	Hunter L	Hunter a	Hunter B						
1	89.63	0.15	5.42						
2	88.10	0.66	6.27						
3	89.98	0.13	4.98						
4	89.53	0.36	4.94						
5	87.76	0.61	5.69						
6	85.83	1.35	6.85						
7	90.79	0.15	4.03						
8	90.67	0.19	4.95						
9	80.10	0.78	4.38						
10	87.23	0.75	6.65						
11	80.70	0.19	3.98						
TOTAL	980.32	5.32	58.14						
MEAN	88.12	0.48	5.29						
S.D.	1.65	0.39	0.99						
RANGE	90.79-85.83	1.35-13	6.85-3.98						
AVER DEV.	1.37	0.31	0.81						
<hr/>									
FIGURE 9									

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5148	Y <sub>xy</sub> , Y	Y <sub>xY</sub> , x	Y <sub>xy</sub> , Y	Lab, L	Lab, a	Lab, b	LCh, L	LCh, C	LCh, h
68-1	60.76	0.34	0.35	82.24	2.32	15.11	82.24	15.28	81.3
68-1	61.89	0.34	0.34	82.82	1.97	14.31	82.85	14.44	82.2
50-2-1	78.39	0.3324	0.3375	90.95	0.68	11.29	90.95	11.31	86.6
50-2-1									
(lint fiber)	21.49	.3155	0.3489	53.48	-8.01	7.97	53.48	11.29	135.2
5148	Hunter L	Hunter a	Hunter B						
68-1	77.94	2.25	13.35						
68-1	78.67	1.92	12.75						
50-2-1	88.53	0.69	10.71						
50-2-1									
(lint fiber)	46.35	-6.35	6.06						

FIGURE 10

5149	Yxy, Y	Yxy, x	Yxy, Y	Lab, L	Lab, a	Lab, b	LCh, L	LCh, C	LCh, h
68-1	65.75	0.3351	0.34	84.86	0.72	11.9	84.86	11.92	86.6
68-1	62.54	.3458	0.3474	83.19	2.14	15.84	83.19	15.98	82.4
68-1	62.56	0.3458	0.3474	83.2	2.14	15.85	83.2	15.99	82.4
8-1	84.72	.3196	0.3278	93.76	0.89	5.87	93.76	5.93	98.6
68-1	64.97	.3316	0.3354	84.46	1.17	9.81	84.46	9.87	83.3
17-2	64.42	.3423	0.3436	84.18	2.26	14.19	84.18	14.36	81
17-3	60.97	.3475	0.3475	82.36	2.74	16.03	82.36	16.26	80.4
17-15-1	64.02	3433	0.3444	83.97	2.34	14.57	83.97	14.75	80.9
21-1	59.32	0.3443	0.3445	81.46	2.64	14.41	81.46	14.64	79.7
21-3	63.64	0.34	0.3409	83.77	2.4	12.89	83.77	13.11	79.5
21-6	67.12	0.3372	0.3394	85.56	1.88	12.15	85.56	12.29	81.3
50-3-1	61.26	0.3502	0.3511	82.51	2.4	17.63	82.51	17.79	82.3
67-1	64.34	0.3434	0.3442	84.13	2.48	14.58	84.13	14.78	80.4
68-1	64.12	0.3442	0.3447	84.02	2.58	14.85	84.02	15.07	80.2
68-2	70.21	0.3428	0.3447	87.09	2.05	15.04	87.09	15.17	82.3
68-3	63.81	0.3457	0.3468	83.86	2.35	15.76	83.86	15.93	81.6
5149	Hunter L	Hunter a	Hunter B						
68-1	81.08	0.71	10.89						
68-1	79.08	2.08	14						
68-1	79.09	2.09	14.02						
8-1	92.04	0.91	5.81						
68-1	80.6	1.15	9.06						
17-2	80.25	2.21	12.75						
17-3	76.08	2.68	14.09						
17-15-1	80.01	2.29	13.05						
21-1	77.01	2.56	12.73						
21-3	79.77	2.35	11.65						
21-6	81.92	1.86	11.14						
50-3-1	78.26	2.33	15.36						
67-1	80.2	2.43	13.07						
68-1	80.07	2.53	13.28						
68-2	83.79	2.04	13.68						
68-3	79.87	2.3	14						

FIGURE 11

5616	Yxy,Y	Yxy,x	Yxy,Y	Lab,L	Lab,a	Lab,b	LCh,L	LCh,C	LCh,h
11-1	72.26	0.3215	0.3254	88.09	1.1	5.06	88.09	5.17	77.8
11-2	58.69	0.3284	0.3335	81.12	0.6	0.36	81.12	0.38	85.9
11-2	52.78	0.3358	0.3335	77.74	3.55	9.22	77.74	9.87	69
11-1	72.03	0.3312	0.3338	87.98	1.72	9.52	87.98	9.67	79.8
11-1	72.34	0.3295	0.332	86.13	1.79	8.64	86.13	8.82	78.4
11-1	71.98	0.3295	0.3313	87.95	2.09	8.39	87.95	8.64	76.1
11-1	73.01	0.3256	0.3305	88.45	0.68	7.51	88.45	7.54	84.9
17-1-2	75.85	0.3274	0.3306	89.78	1.52	7.94	89.78	8.08	79.3
17-3-1	72.6	0.3271	0.3303	88.25	1.48	7.66	88.25	7.8	79.1
17-4-1	69.02	0.3362	0.3377	86.51	1.78	11.37	86.51	11.5	81.2
25-11-1	69.5	0.3364	0.3401	86.75	1.26	12.41	86.75	12.47	84.2
25-28-1	72.21	0.3324	0.3343	88.06	2.09	9.9	88.06	10.11	78.2
25-36-2	70.46	0.3327	0.3353	87.22	1.73	10.22	87.22	10.38	80.5
35-35-1	75.59	0.3268	0.3289	89.66	1.56	7.58	89.66	7.73	78.4
50-12-1	73.13	0.3284	0.3316	88.5	1.46	8.36	88.5	8.48	80.1
KS-11-2	65.33	0.3371	0.3388	84.65	2.07	11.83	84.65	12	80.1
5616	Hunter L	Hunter a	Hunter B						
11-1	85	1.09	4.89						
11-2	76.61	0.58	7.64						
11-2	72.64	3.38	8.22						
11-1	84.87	1.72	8.97						
11-1	85.05	1.79	8.2						
11-1	84.84	2.08	7.98						
11-1	85.44	0.67	7.18						
17-1-2	87.08	1.52	7.62						
17-3-1	85.2	1.48	7.31						
17-4-1	83.07	1.76	10.52						
25-11-1	83.36	1.25	11.43						
25-28-1	84.97	2.08	9.32						
25-36-2	83.94	1.72	9.58						
35-35-1	86.94	1.57	7.29						
50-12-1	86.51	1.46	7.98						
KS-11-2	80.82	2.04	10.81						

FIGURE 12

## **SUBSTITUTE SHEET (RULE 26)**

BC	Yxy, Y	Yxy, x	Yxy, Y	Lab, L	Lab, a	Lab, b	Lch, L	Lch,C	Lch,h
12 Green	33.34	0.3779	0.3717	66.01	4.24	24.18	66.01	24.54	80.1
22 Brown	38.18	0.3778	0.3662	68.15	6.18	23.31	68.15	24.11	75.2
3 Red	24.23	0.4055	0.3728	56.31	10.98	25.52	56.31	27.77	66.9
4 Ivory	46.84	0.3657	0.3599	74.08	4.6	21.13	74.08	21.62	77.8
BC	Hunter L	Hunter L	Hunter a	Hunter B					
12 Green	59.44	3.79	17.92						
22 Brown	61.78	5.62	17.69						
3 Red	49.22	9.42	17.14						
4 Ivory	68.43	4.31	17.02						

FIGURE 13

SUBSTITUTE SHEET (RULE 26)



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## INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification <sup>6</sup> : <b>C12N 15/29, 15/82, 5/10, A01H 5/00</b>		A3	(11) International Publication Number: <b>WO 96/40924</b> (43) International Publication Date: 19 December 1996 (19.12.96)
(21) International Application Number: PCT/US96/09897 (22) International Filing Date: 7 June 1996 (07.06.96)		(81) Designated States: AU, CA, CN, JP, KG, KZ, MX, TJ, TM, TR, US, UZ, European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE).	
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(60) Parent Application or Grant (63) Related by Continuation US Filed on		(88) Date of publication of the international search report: Not furnished (CIP) Not furnished 6 February 1997 (06.02.97)	
(71) Applicant ( <i>for all designated States except US</i> ): CALGENE, INC. [US/US]; 1920 Fifth Street, Davis, CA 95616 (US).			
(72) Inventors; and (75) Inventors/Applicants ( <i>for US only</i> ): McBRIDE, Kevin [US/US]; 1309 Marina Circle, Davis, CA 95616 (US). STALKER, David, M. [US/US]; 2736 Cumberland Place, Davis, CA 95616 (US). PEAR, Julie, R. [US/US]; 818 Douglass Avenue, Davis, CA 95616 (US). PEREZ-GRAU, Luis [ES/US]; 1230 Elk Place, Davis, CA 95616 (US).			
(74) Agents: SCHWEDLER, Carl, J. et al.; Calgene, Inc., 1920 Fifth Street, Davis, CA 95616 (US).			

(54) Title: COTTON FIBER TRANSCRIPTIONAL FACTORS

## (57) Abstract

Novel DNA constructs are provided which may be used as molecular probes or inserted into a plant host to provide for modification of transcription of a DNA sequence of interest during various stages of cotton fiber development. The DNA constructs comprise a cotton fiber transcriptional initiation regulatory region associated with a gene which is expressed in cotton fiber. Also provided is novel cotton having a cotton fiber which has a natural color introduced by the expression in the cotton fiber cell, using such a construct, of pigment synthesis genes. Cotton fiber cells having color produced by genetic engineering and cotton cells comprising melanin and indigo pigments are included.

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## INTERNATIONAL SEARCH REPORT

International Application No

PCT/US 96/09897

A. CLASSIFICATION OF SUBJECT MATTER  
 IPC 6 C12N15/29 C12N15/82 C12N5/10 A01H5/00

According to International Patent Classification (IPC) or to both national classification and IPC

## B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)  
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Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

## C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	WO,A,95 08914 (AGRACETUS) 6 April 1995 see page 8, line 12 - page 9, line 17 see page 18, line 6 - page 20, line 16 see page 36, line 1 - page 38, line 36 see sequence ID nos 4 and 5 ---	1
X	WO,A,94 12014 (AGRACETUS) 9 June 1994 see page 9, line 29 - page 10, line 32 see page 19, line 5 - page 21, line 24 see page 40, line 8 - page 43, line 35 ---	1 -/-

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2 Date of the actual completion of the international search

6 December 1996

Date of mailing of the international search report

20.12.96

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Maddox, A

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International Application No  
PCT/US 96/09897

C(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT		
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X	DATABASE WPI Section Ch, Week 9021 Derwent Publications Ltd., London, GB; Class A35, AN 90-161489 XP002020420 & JP,A,02 104 773 (NAGASE SANGYO KK) , 17 April 1990 see abstract	38
A		13-21, 27-36
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P,X	--- WO,A,95 16783 (CALGENE INC ;MCBRIDE KEVIN E (US); STALKER DAVID M (US)) 22 June 1995 see page 8, line 38 - page 10, line 14	13-18,20
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## INTERNATIONAL SEARCH REPORT

International Application No  
PCT/US 96/09897

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT		
Category	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
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A	WO,A,89 08145 (BIOSOURCE GENETICS CORP) 8 September 1989 see page 5, line 8 - line 18 see page 28, line 17 - line 20 see page 41, line 21 - page 42, line 2 see page 49, line 8 - line 17 ---	16,17
A	WO,A,95 13386 (BIOSOURCE TECH INC) 18 May 1995 see page 23, line 26 - page 24, line 35 ---	16,17
A	WO,A,93 02195 (COMMW SCIENT IND RES ORG) 4 February 1993 see page 1, line 23 - line 29 see page 1, line 36 - page 2, line 2 see page 7, line 1 - line 7 ---	16,17
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